



Department of Mathematics and Computer Science
Data Mining Research Group

Master's Thesis

Towards Increased Adoption of Meta-Learning in AutoML

MLTA: a Meta-Learning Toolbox for AutoML

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Abstract

AutoML promises to transform the current imperative machine learning paradigm into an imperative paradigm, abstracting away machine learning model development. Key to the development of high-performing AutoML is meta-learning, or learning-to-learn, which aims to reuse prior task experience to learn new tasks more quickly. Meta-learning solutions can consist of a variety of components, including similarity measures, characterization methods and the meta-learner itself. However, selecting these components is difficult because we cannot directly assess their performance due to a lack of ground truth values. Therefore, designing and developing meta-learning solutions is challenging and requires implementations to assess them behaviorally. Yet, current AutoML-directed meta-learning research is plagued by issues concerning reproducibility, accessibility, and comprehensibility limiting the adoption of meta-learning in AutoML systems. In this thesis we propose a Meta-Learning Toolbox for AutoML (MLTA) to easily construct, analyze and compare existing and novel meta-learning approaches and components. The toolbox supports AutoML practitioners to adopt meta-learning in AutoML and may also be valuable for meta-learning related research fields including continual learning, multi-task learning and transfer learning. To provide the toolbox with modularity and extensibility properties we devise an AutoML-directed meta-learning framework and formalize the meaning of its main components. Furthermore, we perform a literature analysis on meta-learning components to identify and compare state-of-the-art approaches, incorporating the most promising approaches in MLTA. To showcase MLTA's utility we use it to perform a case study on warm-starting pipeline search. To that end we create a metadataset of nearly 1 million evaluations which we make publicly available. Moreover, we conclude that portfolio building seems to be a promising meta-learning approach for time-restricted settings, but not for warm-starting AutoML systems employing an asynchronous evolutionary algorithm. Our case study suggests that a warm-starting procedure may depend on the subsequent pipeline optimization's algorithm and the set of prior experiences (metadataset). Subsequently, we suggest follow-up work – employing MLTA – to explore the effect of the metadataset, AutoML system, and exploration-exploitation trade off in warm-starting pipeline search.

TODO: update
abstract at end,
mainly with
conclusions

Preface

This thesis concludes my Master's Program Data Science & AI at the Eindhoven University of Technology (TU/e). I would like to thank my supervisor Joaquin Vanschoren and daily supervisor Prabhant Singh for providing their guidance, feedback and support during the execution of this thesis despite their busy schedules. Their insights have helped me to shape and achieve this work. Moreover, I would like to thank the committee members for taking the time to read, assess and discuss my work. In addition to them I would like to thank my co-students for providing valuable insights and discussions. Lastly, I want to thank my family, friends and partner for their unconditional support, motivating me to get the best out of my thesis. Without these people my thesis would not have been possible.

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Chapter 1

Introduction

This introductory chapter intuitively provides the thesis' main concepts and their importance in Section 1.1. Therein, we introduce AutoML, meta-learning, their relation, and provide an example of a meta-learning solution and associated challenges. The main concepts of this thesis are formalized and discussed in-depth in Chapter 2. In Section 1.2 we formulate the research problem and detail the research objectives which are within the overarching research theme of facilitating increased adoption of meta-learning in AutoML. In Section 1.3 we give an overview of the thesis's contents addressing the research aim.

1.1 Motivation and Challenges

Machine learning (ML) concerns models that learn (from data). Currently, most machine learning follows an imperative paradigm, requiring manual specification of model development steps. To make machine learning more accessible, imperative ML can be transformed into declarative ML. In the declarative paradigm, one only needs to specify the desired input and output, without the in-between steps. Going from imperative to declarative ML is where AutoML comes in. Though the goals and outcomes of AutoML are manyfold (see Section 2.1), the main interest is, as the name suggests, in automation of machine learning. Thus, AutoML promises to achieve declarative ML through abstraction of model development.

AutoML has become a vast research field, with popular open-source contributions [57, 40, 51], and multiple companies provide AutoML services [2, 1, 4, 3]. However, for now, most AutoML systems are not end-to-end and are unable to consistently achieve expert-level, human, performance [147]. Moreover, sometimes configuring an AutoML system may be more complex than configuring a ML model. Therefore, more research is necessary to fulfill the AutoML promise of declarative ML. A critical aspect of any AutoML system is efficiently developing ML models because the model development search space is huge. For instance, we can specify the model class, and characterize how it should learn, from differently processed data all with varying contexts. As elucidated, automation of machine learning – especially model development – is far from trivial. It gives rise to the question: “can we learn to efficiently develop learning models?”

One research field in particular is trying to answer that question: meta-learning. Meta-learning is closely connected to the definition of learning: “an increase, through *experience*, of problem-solving ability” [132]. Meta-learning, or learning-to-learn, covers any type of (machine) learning using prior task experience to learn new tasks more quickly [126]. Meta-learning

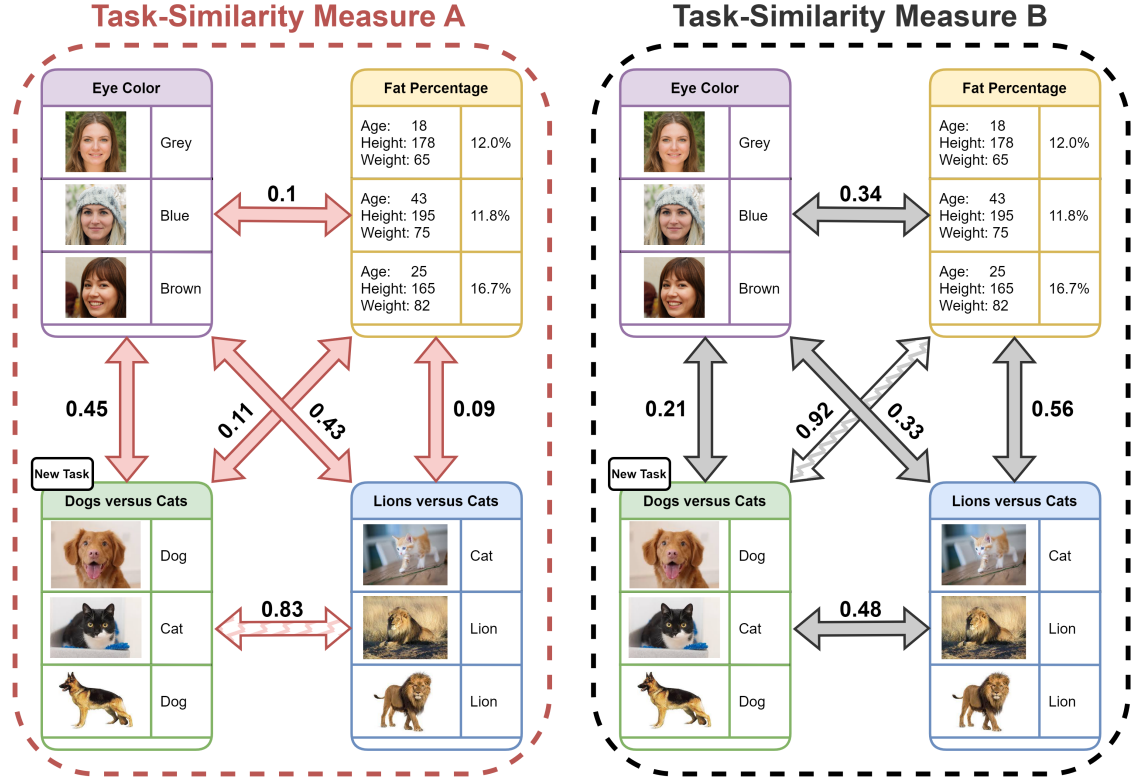


Figure 1.1: An example of a meta-learning application utilizing two *hypothetical* dataset-similarity measures, A and B, on four tasks. This example is created for illustrative purposes, contemporary similarity measures hardly support multiple data modalities. The values near the arrows indicate the *made-up* similarity between the datasets. The humans shown do not exist, they are created with thispersondoesnotexist.com.

covers many different challenges, all with a fundamental notion of reusing previously learned information. Typically, meta-learning research builds upon the assumption that similar tasks are useful as previous experience.

The *assumption*, that previously-seen similar tasks are useful for current learning tasks, is solidified through similar findings in cognitive science [95, 79]. However, with automation in mind, we distantiate from cognitive science by distinguishing between different forms of information: quantitative and qualitative information. Humans can express similarity through qualitative information, but automation is more feasible for quantitative information. Therefore, to enable identifying useful experiences, similarity must be quantified, which is the goal of similarity measures. Likewise, we could employ (quantifiable) characteristics of previous and to-be-solved tasks and their solutions – i.e. dataset and ML model characterizations (see Section 2.2) – to aid learning a novel task. Quantification of task similarity and characterization of tasks and solutions are fundamental to meta-learning, which in turn is suitable to solve challenges in AutoML.

To concretize this thesis' main concepts and their usage we utilize Figure 1.1 illustrating a *simplified and fictional* meta-learning application and the role of similarity therein. Later, in Section 2.2 we provide a more elaborate discussion of meta-learning approaches and their components. Figure 1.1 displays two *hypothetical* task-similarity measures, A and B, assigning

similarity values to pairs of the following four distinct tasks: (1) classifying eye color based on images; (2) predicting the fat percentage using tabular data; (3) classifying dog versus cat images; (4) classifying lion versus cat images. The goal of a task-similarity measure is to quantify the similarity between pairs of tasks, the obtained similarity values can be used for knowledge transfer (meta-learning) applications.

Consider the meta-learning scenario transferring knowledge, say the choice for machine learning algorithm, from previous tasks to a new task. Suppose we want to learn a new task, the dogs-versus-cats classification task. In addition, we have previously solved the three aforementioned tasks: eye color, fat percentage, and lions versus cats. Task-similarity measure A identifies the lion-versus-cat task as the most similar to our current task with a similarity value of 0.83. Instead of learning what algorithm is most suitable for our new task, we could employ a simple meta-learning strategy: use the best-performing algorithm from the most similar task. The ML algorithm from the previously-learned lions-versus-cats task likely transfers well to our new dogs-versus-cats task, in turn accelerating the learning process.

On the other hand, task-similarity measure B identifies the fat percentage task to be the most similar to our dogs-versus-cats task with a similarity value of 0.92. Due to the task's varying data modality (tabular versus image data) we do not expect successful knowledge transfer. Hence, task-similarity measure B likely identifies and transfers an algorithm poorly suited to the new task, consequently yielding inferior performance compared to learning the new task from scratch – a phenomenon referred to as negative transfer [112].

Thus, similarity measures – and other meta-learning components – can influence the performance of meta-learning applications but selecting these components is challenging; we cannot directly measure their performance since there is *no* ground truth. Typical meta-learning applications consist of multiple difficult-to-select components whose performance may be dependent on the application and available prior knowledge. Our simple example already required specification of two interacting components: the similarity measure, and the meta-learner. However, meta-learning approaches may use varying types of components (see Section 2.2.1 and 2.2) further complicating the development of a meta-learning application. Due to dependence on the application and prior knowledge, and the lack of ground truth, we must resort to the implementation of and experimentation with meta-learning approaches.

It may be challenging to design and develop a meta-learning solution, however effectively using meta-learning has several benefits. There exist at least four benefits to using meta-learning, each can be framed as a cost reduction through more efficient learning (requiring less for similar performance). The benefits include reducing costs in experimentation, data collection, training, and missed opportunities. Experimentation costs could be reduced by requiring fewer design options to achieve satiable performance. Data collection – including annotation – costs can be reduced because we may need to collect fewer observations for equal performance. Less data for similar performance may lead to reduced training costs because training time for ML algorithms typically scales with the amount of data. Lastly, for an equal amount of data, effectively using meta-learning may lead to a more suitable model – for instance higher accuracy-related performance – in turn reducing opportunity costs.

1.2 Formulation of the Problem and Objectives

In the previous subsection, we introduced meta-learning solutions and correspondingly established their complexity and explained their potential benefits. Despite an abundance of AutoML systems [85] and meta-learning approaches for AutoML (see Section 2.2), only a limited amount of open-source AutoML systems incorporate meta-learning, examples being Auto-Sklearn [40], Auto-Sklearn 2.0 [38], AlphaD3M [32] and Auto-PyTorch Tabular [146], with 3 out of 4 systems stemming from a single research group. As a main challenge for this thesis we identify the limited adoption of meta-learning for AutoML, and we attribute it to issues concerning reproducibility, accessibility and comprehensibility, which unfortunately reinforce each other.

Existing AutoML-directed meta-learning works tend to be irreproducible and inaccessible because their specifics are not discussed and their implementations not provided. Additional reproducibility issues are caused by approaches' dependence on varying sets of prior experiences – in the form of a metadataset – which is often not made publicly accessible. Furthermore, with varying metadatasets it is challenging to compare meta-learning approaches, limiting comprehension and reducing the impact of any related meta-learning work. Though there is increased research towards meta-learning components, there is no clear overview nor comparison of them, again limiting comprehension and accessibility.

On top of that, many works' implementations for approaches and components are overly tailored to the specifics of their use-case. Subroutines, such as computing dataset characterizations or similarity, are overly dependent on their interpretation, the meta-learning algorithm, and metadatasets' materialization. Due to implementations' overdependence on the use-case convenient re-use is diminished, negatively affecting reproducibility and prototyping pace, successively reducing the number of novel ideas and comparisons between them.

A lack of accessibility is to some extent inherent to complex systems, however, compared with other machine-learning-based research fields the degree of accessibility is limited due to the lack of open-source initiatives providing unified access to implementations. The lack of such initiatives additionally culminates in time-consuming prototyping and experimentation, a problem especially severe for meta-learning components whose performance can only be established through experimentation due to the lack of a ground truth.

To sum up, current AutoML-directed meta-learning research is plagued by issues concerning reproducibility, accessibility and comprehensibility, stemming from the lack of standardized implementations, difficulty in creating prototypes and comparisons, and the absence of an AutoML-focused meta-learning framework. The lack of reproducibility, accessibility and comprehension limits the adoption of meta-learning for AutoML. Striving towards increased adoption of meta-learning in AutoML systems, we propose to alleviate the aforementioned issues by introducing a Meta-Learning Toolbox for AutoML (MLTA), focusing on tabular data for supervised classification. MLTA provides a modularized and extensible framework for AutoML-directed meta-learning solutions, standardizing development through its framework, providing unified access to common components, facilitating comparisons and promoting code-reuse through its modularization. The framework and toolbox components are designed agnostically with respect to the AutoML-related task it solves, e.g. warm-starting search, performance prediction or training time estimation. Lastly, we hope that through promoting a pursuit of standardized practices we may inspire the community to arrive at a consensus for comparing meta-learning solutions in the form of a benchmark.

The aforementioned challenge – increased adoption of meta-learning in AutoML – sets the stage for this thesis, but it does not allow assessing its success. Therefore we define a more concrete research objective with respect to MLTA:

1. create a toolbox to easily construct, analyze and compare existing and novel meta-learning approaches and components.

In a broader sense, we thereby aim to *partially* mitigate reproducibility, accessibility and comprehensibility issues (with meta-learning research) to support AutoML practitioners to adopt meta-learning in AutoML. This thesis is most useful for research(ers) in the areas of AutoML, meta-learning and related research areas.

In Section 2.2 we explain the relation of multi-task learning, transfer learning, NAS and continual learning to meta-learning. These, and additional, related research fields could employ components which are present in meta-learning systems, making them potential beneficiaries of our toolbox. To support that claim we provide an overview of the usage of similarity measures in the aforementioned research fields in Appendix A. Furthermore, through potential accelerated research towards AutoML systems, there may be a benefit for their users. These can be expert and non-expert ML enthusiasts from companies or completely different research fields.

The proposed toolbox distinguishes itself from other meta-learning toolboxes by its focus on AutoML systems for supervised tasks on tabular datasets. In contrast, current meta-learning toolboxes such as learn2learn [13], NASLib [113] TorchMeta [28] and BOML [78] are mostly directed towards gradient-based methods in NAS, meta-reinforcement learning, and few-shot learning. MLTA differs from works providing meta-learning components for tabular datasets, such as MFE [9], by its broadness focusing on all main components of a AutoML-related meta-learning application. MLTA extends MFE’s dataset characterization functionality by additionally including more recent, and potentially higher-performing, dataset characterization methods and corresponding similarity measures. Apart from MLTA’s divergent focus, MLTA distinguishes itself from other approaches by explicitly incorporating prior experiences (ML pipelines) in its meta-learning framework.

This thesis shares great similarity with GAMA [47], which aims to support the development of novel ideas for AutoML systems and its components by providing a modularized framework. Like GAMA, our work intends to support AutoML research, but it distinguishes from GAMA by its focus on meta-learning for AutoML – a feature currently not incorporated in GAMA. The need for our toolbox is strengthened by the success of related works such as GAMA, MFE, learn2learn and NASLib.

1.3 Thesis Outline

The remainder of this thesis is structured as follows. In Chapter 2 we explore our problem setting in-depth, detailing AutoML and exploring the state-of-the-art for AutoML-associated meta-learning. Moreover, in Chapter 2, we provide a structured overview of existing approaches for typical meta-learning components, which serves as a basis for Chapter 3 introducing "MLTA", the meta-learning toolbox. In Chapter 3, we compare the state-of-the-art presented in Chapter 2 to decide which components should be provided by the toolbox. In Chapter 4 we showcase the toolbox achieves the research objective by performing a case study – employing MLTA – on warm-starting pipeline search. In Chapter 5 we summarize key take-aways and reflect on our contributions by contextualizing them and assessing their implications.

Chapter 2

Preliminaries

In this chapter we introduce the main preliminaries of this thesis: AutoML, associated meta-learning solutions and their components. In their respective subsections we provide formulations, where appropriate a formalization, and explain the relation to the thesis' goals. This chapter additionally serves as an overview of state-of-the-art AutoML-related meta-learning works, explaining necessary details for their suitability comparison in Chapter 3.

2.1 AutoML

2.1.1 Problem Setting and Toolbox Scope

In Section 1.1 we already established that AutoML promises to make machine learning declarative, instead of imperative. In [147] it is explained that AutoML aims to make design decisions in a data-driven, objective, and automated way. The authors exhibit the vast size of the research field by characterizing the numerous challenges in AutoML, they include:

"automating data collection and experiment design; automating data cleanup and missing data imputation; automating feature selection and transformation; automating model discovery, criticism, and explanation; automating the allocation of computational resources; automating hyperparameter optimization; automating inference; and automating model monitoring and anomaly detection."

The field of AutoML is vast and diverse, as apparent from the many distinct challenges, therefore we need to scope the toolbox by restricting ourselves to AutoML for:

1. Model development,
2. on tabular datasets,
3. in a supervised learning setting,
4. performing classification tasks,
5. considering non-deep-learning ML pipelines (see 2.1.3).

We introduce restriction (5), to preclude deep-learning models and corresponding pipelines, to avoid overlap with the research field of Neural Architecture Search (NAS). It does however not

restrict the toolbox' usage in NAS works; the toolbox could support works using both classical and NN-based ML models like Auto-PyTorch Tabular [146].

Restriction (2) concerns the focus on tabular data, whose definition we adopt from [20], namely: a heterogeneous dataset with a fixed number of features that are either continuous or categorical, typically presented in a table with data points as rows and features as columns. Restricting ourselves to tabular data strengthens the focus on AutoML and avoids overlap with NAS, because tabular data's properties favor more traditional machine learning methods over deep learning [20].

Note that the terminology for restriction (1), "model development", does not occur in the aforementioned collection of AutoML challenges, because it encompasses multiple challenges including: automating feature selection and transformation; automating model discovery; automating hyperparameter optimization and automating the allocation of computational resources. Such challenges could be partially solved by employing meta-learning systems, which for instance carry out performance predictions or training time estimation. Thus, meta-learning solutions and components facilitated by MLTA can help achieve high-performing AutoML, desirable for a variety of reasons. An AutoML system would enable non-experts to build ML applications [147]. Moreover, in the pursuit of increasing accessibility and democratization, we obtain tools and practices for more systematic and efficient machine learning [55]. An example of such a tool is Hyperband [73], for the challenge of hyperparameter optimization, whose success is demonstrated through its incorporation in popular libraries such as Keras [25].

2.1.2 Problem Formulation

In AutoML we search for solutions, termed configurations, in a configuration search space. AutoML systems vary by their search method and search space, popular frameworks include Autogluon [35], auto-sklearn [40], auto-sklearn 2.0 [37], FLAML [129], GAMA [47], H2OAutoML [71], AutoWeka [51] and TPOT [98]. Thus, the definition of a configuration differs across its usage, but typically a configuration is a choice in the model development search space. A configuration in the full model development search space thus includes any aspect defining a part of the model as a whole, it can include options for hyperparameters, algorithms, pipelines, architectures, and so forth. A main task of AutoML systems is to find an adequate configuration, typically by searching the configuration space, which is challenging due to multiple factors [39]. First, evaluations can be, and typically are, expensive. This issue worsens over time, with developments of (extremely) complex models, such as Megatron-Turing NLG containing 530 billion parameters [118]. Moreover, datasets are getting larger, increasing evaluation time even further. Second, the configuration space is often complex and high-dimensional. Furthermore, it may not be clear which dimensions of the configuration are important. Not only is choosing an adequate configuration challenging, but it is also important. There are numerous findings indicating the effect of configuration choice on performance. For instance, in hyperparameter tuning, performance can be improved over the default settings [83, 97, 115]. Moreover, weaker well-tuned models can outperform stronger, poorly-tuned, models [84]. Additionally, findings suggest performance depends on configuration choices for the model [97], model and hyperparameters [123], pre-processing techniques [8, 108], and architecture [80, 139].

The exact materialization of what constitutes a configuration, or its search space, depends on the AutoML formulation that is chosen. Some works, like H2OAutoML [71], employ the Combined Algorithm Selection and Hyperparameter Optimization [123] formulation commonly

termed CASH. Others, like TPOT [98], additionally include feature engineering steps, and some approaches extend this by including data cleaning steps such as AutoWeka-MCPS [114]. In this thesis, we want to focus on a formulation of AutoML allowing us to provide the most general insights about meta-learning for AutoML. We argue for the formulation of AutoML that searches for preprocessing steps performing feature engineering, algorithm selection and hyperparameters for each of those, an approach taken for instance by the AutoML frameworks TPOT [98] and GAMA [47]. By including preparatory steps in our formulation we align better with the goal of true automation of ML, because it is reported that practitioners spend the majority of their time in such preparatory steps [104]. This formulation is similar to CASH, but additionally includes preprocessing steps performing feature engineering. On the other hand it searches for pipelines, like approaches additionally taking into account data preparation steps. However, in contrast to those approaches, our formulation does not include data preparation steps because information transferability is less straightforward for them due to their inherent dependence on the dataset.

2.1.3 Problem Formalization

Our formalization of the AutoML problem, presented in Definition 1, follows that of [46] adapting the interpretation of [147]. It generalizes previous formulations and interprets AutoML as a pipeline creation problem with the objective to minimize the empirical risk of a pipeline according to some evaluation procedure. Interpreting a pipeline as a DAG, the formalization decomposes AutoML by separately considering the pipeline structure and its materialization in the form of algorithms and hyperparameters.

Definition 1 (Pipeline Creation Problem). *Let a set of algorithms \mathcal{A} with an according domain of hyperparameters $\Lambda_{(.)}$, a set of valid pipeline structures G and dataset \mathcal{D} be given. The pipeline creation problem consists of finding a pipeline structure in combination with a joint algorithm and hyperparameter selection that minimizes the loss*

$$(g, \mathbf{A}, \boldsymbol{\lambda})^* \in \underset{g \in G, \mathbf{A} \in \mathcal{A}^{|g|}, \boldsymbol{\lambda} \in \Lambda}{\operatorname{argmin}} \mathcal{R}(\mathcal{P}_{g, \mathbf{A}, \boldsymbol{\lambda}}, \mathcal{D}) \quad (2.1)$$

within a given resource budget B .

where:

- $g \in G$ is a graph from the set of all valid graphs,
- $\mathbf{A} \in \mathcal{A}^{|g|}$ is a vector specifying for each node in graph g the algorithm from the set of algorithms \mathcal{A} ,
- $\boldsymbol{\lambda} \in \Lambda$ is a vector specifying the hyperparameter configuration of each algorithm from the set of all possible configurations Λ ,
- B is a resource budget, for instance in time or iterations.

Note that the AutoML formulation, through its interpretation as a DAG-based pipeline creation problem, is rather broad. Therefore, to align the formalization with the focus of our work we impose one further restriction for this thesis. For the search space of the pipeline creation problem we explicitly do not consider algorithms in \mathcal{A} performing data preparation

steps, e.g. those performing formatting, encoding and imputation. This need not imply pipelines lack these steps, they may for instance be fixed or rule-based. We simply state that we exclude data preparation algorithms from the search space because, with meta-learning in mind, they are difficult to transfer due to their inherent dependence on the dataset. We impose no further restrictions on G , B , Λ and \mathcal{A} nor the evaluation procedure estimating \mathcal{R} , because these specifications are relevant for a particular AutoML system or use-case, not for the meta-learning toolbox introduced in this thesis.

2.2 Meta-Learning

In Section 1.1 we discussed meta-learning, or learning-to-learn, and its relation to similarity measures. A brief recap, meta-learning is “the science of systematically observing how different machine learning approaches perform on a wide range of learning tasks, and then learning from this experience, or meta-data, to learn new tasks much faster than otherwise possible. Additionally, meta-learning approaches typically assume similar tasks to be useful previous experiences. Identifying the main concepts in the definition of meta-learning and its assumptions yields a list of main components for a meta-learning system: a set of prior experiences (metadataset), machine learning approaches (configurations), a learning task (dataset) and learning the new task (meta-learner). In the forthcoming Subsections we discuss methods related to these main components in more depth, except for the metadataset because we are unaware of any works explicitly investigating it. In addition to that, we provide an overview of the state-of-the-art. This preliminary Section is most related to a meta-learning survey [126], but distinguishes itself by including more recent approaches and the explicit focus on and distinction between AutoML-related meta-learning components. Before exploring the main components of meta-learning systems we outline the field of meta-learning and its related fields.

The broad definition of meta-learning allows it to include lots of different approaches or even entire research fields such as transfer learning or Neural Architecture Search (NAS). Though current meta-learning research is mostly focused on applications in deep learning [53], the research field is far broader. According to [126] meta-learning approaches are typically part of one of three categories, they either learn from model evaluations, task properties, or prior models. The latter refers to transfer learning, which concerns approaches that account for differences in two distributions [63], commonly termed a source and task distribution. It accounts for these differences using knowledge transfer techniques. Transfer learning can be seen as a form of meta-learning where previously learned models are transferred.

Besides transfer learning and NAS, the field of meta-learning is related to multi-task learning and continual learning. Multi-task learning (MTL) is an approach using information in the training signals of related tasks; utilizing a shared task representation [21]. Like meta-learning, MTL aims to use information learned from other tasks. MTL uses task information by jointly learning tasks, while meta-learning learns tasks separately. Hence, in MTL you learn from a task by learning it directly, while in meta-learning you learn from an already completed task. Continual learning considers a setup to learn continually from an infinite stream of data trying to gradually extend knowledge and use it for future learning [27]. From the definition of continual learning, we point out the connection to meta-learning and multi-task learning. As with meta-learning, continual learning aims to reuse learned information. As with multi-task learning, continual learning learns from more than one task, though not simultaneously. Distinguishing it from MTL and meta-learning, continual learning not only aims to learn from

other tasks, but to keep on learning.

2.2.1 Characterization Methods and Similarity-Measures

In this subsection we discuss preliminary work on configuration characterizations, dataset characterizations and dataset similarity measures, which are common meta-learning components typically interoperating with a meta-learner (see Section 2.2.2). According to the Cambridge Dictionary a characterization is "a description of the most typical or important characteristics of someone or something" while Oxford English Dictionary defines it as "a description of the distinctive nature or features of someone or something". Hence, a dataset characterization is a description of the most important or typical characteristics of a dataset or its distinctive nature. Comparably, a configuration characterization describes the distinctive nature or most important or typical characteristics of a configuration. Though similar to a dataset characterization, a dataset similarity measure solely quantifies the similarity or dissimilarity between datasets. In this thesis, we discriminate between dataset characterizations and similarity measures because their materialization and usage vary despite many works simultaneously providing solutions for both.

Typically a dataset characterization takes the form of a vector (or matrix), while similarity measures represent the (dis)similarity between datasets using a scalar value, i.e. a similarity value. Their usage varies with some meta-learners requiring the characteristics of a dataset, while others only employ its relation to other datasets (see Section 2.2.2). Distinguishing between properties, a dataset characterization method can operate in isolation of other datasets while a similarity measure cannot.

As mentioned, many works provide solutions for both a dataset characterization method and a similarity measure. After all, a dataset characterization method providing fixed-size vectorized representations of datasets allows to be extended to a similarity measure by computing the similarity between characterizations. Please note the ambiguity of the term "*dataset characterization*" — either referring to a dataset's distinctive nature *or* its characteristics — captures its possible extension well. However, a characterization method is not per se extensible because a characterization's dimensionality may depend on properties of the dataset it characterizes.

A characterization method extended to be a similarity measure benefits from a key property of a characterization method: a characterization exists in isolation of other datasets. This property implies a characterization-based similarity measure can be computed efficiently with respect to the number of datasets; to compute all N^2 pair-wise similarity values between N datasets we only need to execute a characterization method N times. Contrarily, a similarity measure typically needs to be executed N^2 times to compute all N^2 pair-wise similarity values.

To measure the (dis)similarity between two equally-sized vector-based characterizations we can chose among multiple approaches, including the cosine similarity or Euclidean distance. A similarity measure employing the former assumes the difference in angles captures the similarity well, while the latter assumes distances do. It is unclear which approach is most appropriate, the authors of Task2Vec [5] show the dataset characterization's norm encodes the datasets' difficulty, while another work [89] shows the angles between their dataset characterizations are preserved when varying meaningless changes in training factors. These findings highlight more research is needed to choose approaches for measuring the (dis)similarity between characterizations in a well-substantiated manner.

To be precise in our terminology and transparent in our classification of approaches we

provide formal definitions of the previously discussed concepts.

Definition 2 (Dataset Characterization Method). *A dataset characterization method $d : \mathcal{D} \mapsto \mathcal{C}_{\mathcal{D}} \in \mathbb{R}^{n \times m}$ with $n \in \mathbb{Z}^+$, $m \in \mathbb{Z}$ maps a dataset to a characterization $\mathcal{C}_{\mathcal{D}}$ representing \mathcal{D} 's main characteristics. Where a dataset $\mathcal{D} = \{(\mathbf{x}_i, \mathbf{y}_i)\}_{i=1}^N$ consists of feature vectors $\mathbf{x}_i \in \mathbb{R}^d$ having $d \in \mathbb{Z}^+$ features and target vectors $\mathbf{y}_i \in \mathbb{R}^k$ having $k \in \mathbb{Z}^+$ dimensions.*

Note that typically $k = 1$ for a target vector $\mathbf{y}_i \in \mathbb{R}^k$ having $k \in \mathbb{Z}^+$ dimensions but not always, e.g. with multi-label classification $k > 1$. Note we do not include a property for a characterization specifying it uniquely identifies a dataset to allow for a broad overview. Whether this encoding-like property would be preferable for dataset characterization methods is unknown.

Definition 3 (Pipeline Configuration Characterization Method). *A pipeline configuration characterization method $c : (g, \mathbf{A}, \boldsymbol{\lambda}) \mapsto \mathcal{C}_c \in (\mathbb{R} \cup B^*)^{n \times m}$ with $n \in \mathbb{Z}^+$, $m \in \mathbb{Z}$ maps a pipeline configuration to a characterization \mathcal{C}_c representing its main characteristics. A pipeline configuration $(g, \mathbf{A}, \boldsymbol{\lambda})$ is a tuple consisting of a pipeline structure, corresponding algorithms and hyperparameters with $g \in G$ is a graph from the set of all valid graphs, $\mathbf{A} \in \mathcal{A}^{|g|}$ is a vector specifying for each node in graph g the algorithm from the set of algorithms \mathcal{A} , and $\boldsymbol{\lambda} \in \Lambda$ is a vector specifying the hyperparameter configuration of each algorithm from the set of all possible hyperparameter configurations Λ . Let $B = \{a, b, \dots, y, z\} \cup \{0, 1, \dots, 8, 9\}$ be the alphabet consisting of the union of all letters in the Latin-script alphabet and all single digits in the decimal system. Let a string $s \in B^*$ be a sequence of characters in B where B^* denotes the Kleene star on alphabet B .*

In Definition 3 we do not specify which configuration characteristics are captured by the pipeline configuration method. The lack of its specification permits methods only capturing part of the configuration, e.g. those neglecting the hyperparameter configuration or the pipeline structure. This imprecision is deliberate, for it allows a broad spectrum of methods to be consistent with the definition. Moreover, we are deliberately ambiguous with respect to the realization of characterization, it can be a vector or matrix consisting of real-valued numbers and/or strings (a sequence of characters). The broad definition is necessary, because currently only few such methods exist (see 2.2.1.3) and we must avoid steering development in a particular direction by tailoring the definition to a specific method. A pipeline configuration characterization method is akin a dataset characterization method, but for configurations. Therefore, we postulate the possibility to have configuration similarity measures, extending their corresponding characterizations. We are however not aware of any such works and will thus not consider them in this preliminary Chapter. We mention its possibility because we envision potential usage in AutoML systems, for instance a configuration similarity measure could be employed in enforcing diversity in warm-starting evolutionary search frameworks like TPOT [98] or GAMA [47].

Definition 4 (Dataset-Similarity Measure). *A dataset-similarity measure $s : \mathcal{D}_1 \times \mathcal{D}_2 \mapsto \mathbb{R}$, is a mapping from two datasets \mathcal{D}_1 and \mathcal{D}_2 to a real number indicative of the (dis)similarity between the datasets. Where the two datasets are given as $\mathcal{D}_1 = \{(\mathbf{x}_{1,i}, \mathbf{y}_{1,i})\}_{i=1}^{N_1}$ with feature vectors $\mathbf{x}_{1,\cdot} \in \mathbb{R}^{d_1}$ having $d_1 \in \mathbb{Z}^+$ features and target vectors $\mathbf{y}_{1,\cdot} \in \mathbb{R}^{k_1}$ having $k_1 \in \mathbb{Z}^+$ dimensions, and $\mathcal{D}_2 = \{(\mathbf{x}_{2,i}, \mathbf{y}_{2,i})\}_{i=1}^{N_2}$ with feature vectors $\mathbf{x}_{2,\cdot} \in \mathbb{R}^{d_2}$ having $d_2 \in \mathbb{Z}^+$ features and target vectors $\mathbf{y}_{2,\cdot} \in \mathbb{R}^{k_2}$ having $k_2 \in \mathbb{Z}^+$ dimensions.*

Our formalization of a dataset-similarity measure (Definition 4) allows for datasets with varying number of instances, features and target dimensionality. The subsequent overview also includes measures with additional restrictions, which are indicated if present. Observe that in our definition of a dataset-similarity measure we restrict ourselves to pair-wise measures. We claim most interesting applications of similarity measures consider pair-wise similarities, accordingly in this thesis we only consider pair-wise similarity measures. Thus, a dataset similarity measure quantifies the similarity or dissimilarity between exactly two datasets.

In the overview of dataset similarity measures we focus on measures relying on both targets and features, because we have restricted ourselves to a supervised setting (see Restriction 2). It is possible to quantify the similarity between datasets with and without using targets, but there are considerable differences between the power and applicability of these measures as showcased in [5, 11]. For instance, measures incorporating targets are suited to distinguish between two datasets with equal features but varying targets. In contrast, measures which do not take into account targets cannot distinguish between different tasks using the same features. On the other hand, those measures' non-reliance on targets can be a strength, it makes them suitable for unsupervised settings. In this preliminary Chapter, we do not discuss similarity measures solely relying on features because research towards them is limited and due to our supervised setting.

Transferability measures Different from, but very much related to similarity measures, are transferability measures. Transferability measures, also known as transferability metrics, quantify or predict the amount of transferability between a source and a target task [6]. Similarity and transferability measures differ on two main aspects, their broadness and their availability of ground truth information. Transferability measures are less broadly applicable than task-similarity measures because their inputs (may) differ. Typically, transferability measures include source models in their input, whereas similarity measures take *only* datasets as input. Moreover, some transferability measures make additional assumptions on their input. Often the source models are assumed to be neural networks, because they allow for the transferability of their structure and parameters. Transferability measures are model-oriented and more specific due to additional restrictions on the input. This makes transferability measures suited for specific use cases like transferring only certain parts of a neural network [106], or ranking checkpoints to transfer from [75]. Transferability measures only relying on data to characterize the dataset can be framed as dataset-similarity measures, yet not all similarity measures are transferability measures. Unlike similarity measures, the development of transferability measures is guided by ground truth. That is, we can compare transferability measures on their performance in transfer learning. For similarity measures there is no ground truth, one can only evaluate their use for downstream applications such as a meta-learning application. Transferability measures can be used in meta-learning applications, but these are typically not oriented towards tabular AutoML. Therefore, we do not provide an elaborate overview of transferability measures. However, due to their resemblance with similarity measures, possibly influencing their development, we give a brief overview. Popular transferability measures include LEEP [92], LogMe [142] and GBC [100]. These methods measure how well a model transfers to a target task or dataset. To estimate the transferability they use the source model to embed the target data. Other transferability measures include [14, 120, 54, 7, 75, 121], for a more elaborate discussion on transferability metrics consider [6, 122].

Up until now we have introduced configuration characterizations, dataset characterizations and similarity measures, what follows are overviews of them (see Sections 2.2.1.3, 2.2.1.2 and 2.2.1.1). In these overviews we identify the state-of-the-art and provide corresponding explanations introducing their main aspects. Thus, assessing their aspects to determine their suitability for meta-learning is not part of this preliminary chapter, rather, this is carried out in Chapter 3 (detailing the meta-learning toolbox). One particular approach is excluded from the overview which does not conform to the provided Definitions 2, 3 and 4 of a configuration characterization method, nor a characterization method or similarity measure. The approach [33] computes a language-based embedding of the metadata as a representation of the task. We exclude the approach because the manifestation of metadata is greatly dependent on human interpretation. Note that datasets are also products of human interpretation, but they tend to be more objective, specific, complete, consistent, and less language dependent.

2.2.1.1 Dataset Similarity Measures

OTDD Optimal transport dataset distances (OTDD) [11] can be considered a model-agnostic task-dissimilarity measure. Thus, computing the OTDD does not involve training a model. The OTDD is a hybrid Euclidean-Wasserstein distance, it possesses the properties of a distance metric. Hence, the output of OTDD is inversely related to the similarity between two tasks. OTDD treats the distance between two samples as a sum of their feature-based and label-based distance (see Figure 2.1). The method assumes to have access to a metric to compute the feature-based distance. For the label-based distance, OTDD models labels as distributions over feature vectors, therefore the label sets can be completely disjoint. The distance between the label distributions is determined by computing the statistical divergence between their feature-based distributions. Optimal transport is used as the measure for statistical divergence. Then, the distance between the two datasets can be determined by computing the optimal coupling between its samples. Computing both the feature and label-based distances is computationally expensive, therefore the authors propose to approximate the label distribution as a Gaussian distribution. Although this computational trick speeds up computation, OTDD still scales cubically in the number of features, and quadratically in the number of samples. Despite OTDD’s computational burden, the paper does not compare it to baselines to justify the computational complexity.

2.2.1.2 Dataset Characterization Methods

Meta-Feature-Based Characterization Perhaps the most straightforward dataset characterization method is to compute meta-features: an approach similar to the featurization of datasets, but featurizing datasets themselves instead of samples therein to capture dataset’s main characteristics. Thus featurization is employed at one higher level of abstraction (dataset over sample therein) and therefore the featurized representation of the dataset is termed meta-features. More formally, a meta-feature f is a function $f : \mathcal{D} \mapsto \mathbb{R}^k$ that upon application to a dataset \mathcal{D} returns a set of k values characterizing the dataset [110]. Moreover, these meta-features are often hand-crafted to be predictive of the performance of machine learning models applied to dataset \mathcal{D} .

We reuse the terminology and formalization from the authors of [110], whom distinguish between meta-features and characterization measures because a characterization measure $m : \mathcal{D} \mapsto \mathcal{R}^{k'}$ results in k' values, with k' potentially depending on the dataset itself. Whereas

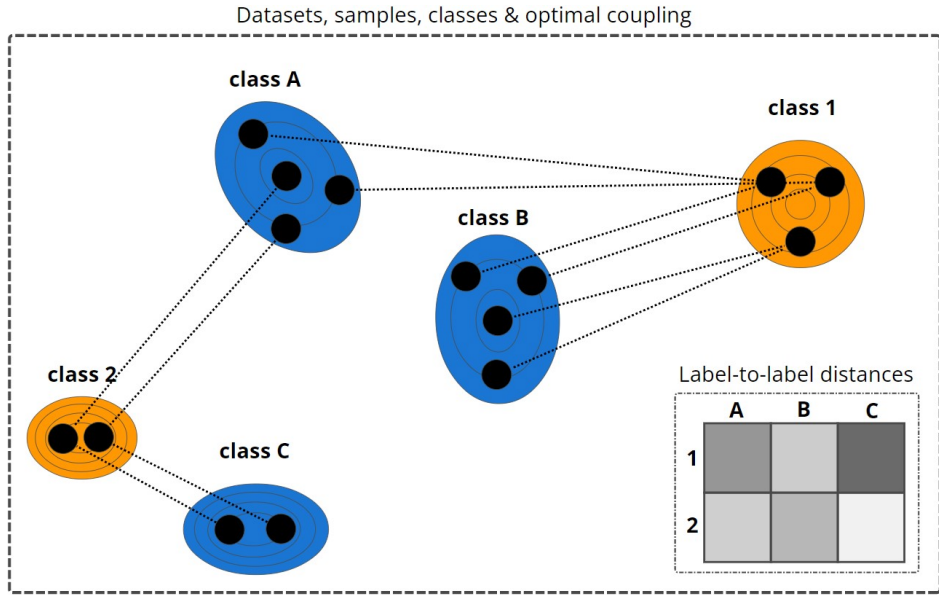


Figure 2.1: OTDD [11] example depicting two *made-up* 2-dimensional datasets with their classes: dataset blue with classes A, B and C; and dataset orange with classes 1 and 2. Data points are depicted as black dots, within their Gaussian-approximated label distribution, whose color depicts the dataset they belong to. The lower right corner depicts the inter-dataset label-to-label distances (darker is larger) which are computed using the Gaussian approximations, which together with a ground metric on features allows to compute the optimal coupling between the datasets' data points (grey dashed lines). Observe dataset blue contains 10 data points while dataset orange contains half of that. Thus, if the mass from one blue point only transfer to exactly one orange point (which happens in the example but need not be the case) there must be 2 blue data points connected to each orange data point. Also note the datasets' label sets are completely disjoint, a key property of OTDD available due to labels being represented by the Gaussians approximating their feature-based distributions.

using meta-features as a dataset characterization requires fixed dimensionality (see Definition 2). Therefore, we must map each characterization measure to a fixed-dimensional meta-feature using a summarization function $\sigma : \mathbb{R}^{k'} \mapsto \mathbb{R}^k$; frequently used summarization functions include the mean, minimum, maximum, skewness and kurtosis. Then, a meta-feature-based dataset characterization is a collection of meta-features $f(\mathcal{D}) = \sigma(m(\mathcal{D}, h_m), h_s)$ with h_m and h_s being hyperparameters for m and σ respectively.

The selection of meta-features is a notoriously difficult task; there are many meta-features for many different tasks whose selection may be context dependent. Within our supervised classification setting, meta-features can be categorized in different groups: simple features easily extracted and general in nature; statistical features capturing statistical properties; information-theoretic features capturing the amount of information and complexity; model-based features derived from properties of models; landmarking features capturing performance on simple and fast learners; and other features. For an extensive discussion of possible meta-features for classification tasks consider [110, 9].

Popular and typically well-performing collections of meta-features for supervised classification settings include the ones proposed by Feurer et al. [43] and Wistuba et al. [134], whose specific meta-features are listed in Appendix C. The former contains simple, information-theoretic, statistical, PCA-based and landmarking meta-features which together with specified

summarization functions result in a dataset characterization containing 46 values. The latter is composed of statistical and information-theoretic meta-features which together with specified summarization functions yield a dataset characterization composed of 22 values. The latter collection of meta-features and summarization functions is a subset of the former, and thus requires fewer computational resources. Perhaps a counter-intuitive finding, the subset outperforms the superset in [58].

Task2Vec Task2Vec [5] characterizes a task in a behavioral way, by observing a neural network’s parameter sensitivity to the data. Given a dataset consisting of images, Task2Vec computes a fixed-size embedding of the dataset using a pre-trained neural network, which is referred to as a probe network. Prior to computing a task embedding, the classification layer of the probe network is re-trained on the given task. The embedding computation builds on the behavioral intuition that neural networks’ activations of the data characterize the data itself. A dataset is passed through the probe network to estimate (the diagonal elements of) the Fisher Information Matrix corresponding to the probe network’s filter parameters. Collecting these estimates in a vector yields the task embedding. Any task’s embedding is of fixed size, because Task2Vec uses the same probe network (architecture) for all embeddings. The authors find that the norm of the embedding encodes the task’s complexity. There are two task-similarity measures based on the Task2Vec embedding, the symmetric Task2Vec distance, and the asymmetric Task2Vec “distance”. Both distances take two Task2Vec embeddings as input and produce a scalar similarity value. However, the asymmetric Task2Vec distance takes the complexity of the first task into account. Therefore, it is asymmetric and not a true distance metric. Moreover, the asymmetric distance introduces hyperparameter α , weighting the influence of complexity. Due to the inclusion of complexity and α , the asymmetric distance can be negative.

Dataset2Vec Another dataset characterization method employing a pre-trained model is Dataset2Vec [58]. Dataset2Vec learns a meta-feature extractor by training a neural network. The last layer of the neural network represents the learned meta-features, together they form a fixed-size embedding of the dataset. The paper itself only presents the embedding method, but does not discuss how to compute similarity or dissimilarity from it. However, applying the cosine similarity on the embeddings is a natural way to compute the similarity value. Neural networks require input consisting of equal size and therefore Dataset2Vec interprets datasets as hierarchical sets, specifically single predictor-target pairs, to achieve schema-agnosticism. To learn a meaningful embedding, the neural network is trained on a novel task with abundant data: The authors propose a patch identification task, where the neural network must predict whether two patches are sampled from one dataset or two distinct datasets. A patch consists of a subset of instances, features, and targets from a given dataset. In performing the patch identification task, Dataset2Vec learns features to discriminate the datasets, in turn yielding the meta-feature extractor. Instead of explicitly learning meta-features for each pair of datasets, the authors pre-trained the network on the patch identification task for a large collection of classification datasets. This pre-trained network extracts the general, previously learned, meta-features that compose the fixed-size task embedding by averaging the meta-features over the number of batches specified.

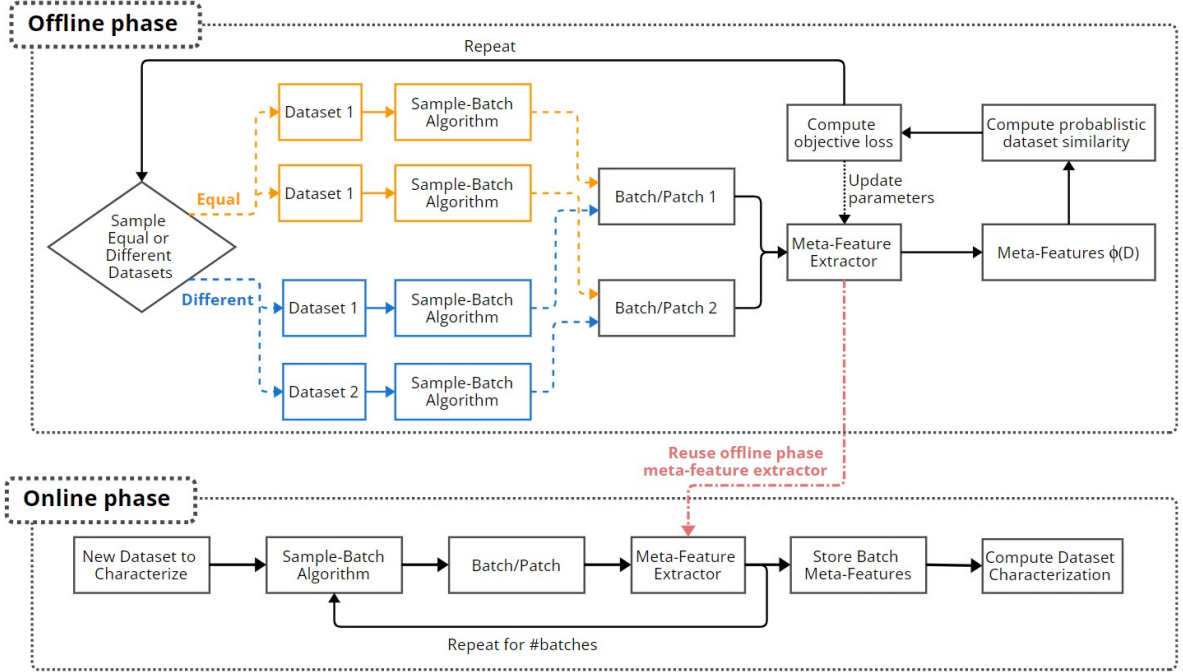


Figure 2.2: Illustration of the Dataset2Vec [58] framework to characterize datasets. The top part of the figure depicts the offline phase, in which the meta-feature extractor is trained. The bottom part depicts the online phase in which a new dataset is characterized using the pre-trained extractor. Solid arrow lines indicate the always-executed flow of steps within a phase, and dashed arrows indicate possible execution flow. The red semi-dashed arrow highlights the interaction between the offline and online phase.

DIDA DIDA [26] is an approach for learning a dataset characterization using distribution-based invariant neural networks. DIDA uses the coordinates in the latent space of the last neural network layer as the task embedding. Therefore, DIDA yields a fixed-dimensional vector consisting of learned meta-features. The authors propose a distribution-based invariant layer as the first layer, or the first two layers, of the neural network. The invariant layer is built on top of aggregated interaction functionals, themselves employing a four-dimensional aggregator, aggregating all feature and label values. By using the invariant layer as the first layer in the proposed DIDA component the authors achieve invariance with respect to labels and features, but at high computational expense. To reduce the computational burden of the invariant layer, the authors propose to only extract and aggregate information from the neighborhood of samples. To learn the meta-features a similar approach to that of Dataset2Vec is used. A Siamese architecture consisting of DIDA components is trained on a patch identification task. The Siamese network is optimized through a loss defined using the distance between the embeddings of two DIDA components, each taking in one dataset. Then, after training, the last layer of the DIDA component represents the learned meta-features. Thus, using DIDA to compute a task-similarity value involves training a neural network, with a considerable amount of parameters. The paper claims DIDA outperforms Dataset2Vec, but the evaluation to support this claim is weak. The only comparison to Dataset2Vec is in the patch identification task, which is not the goal of the learned meta-features, merely a tool to learn them. The paper lacks a comparison to DataSet2Vec in a downstream application, although it does evaluate

DIDA applied to a downstream application.

WTE The Wasserstein Task Embedding (WTE) [77] is an optimal-transport-based dataset embedding method and the Euclidean distance between the embeddings produces an associated dataset similarity value. The WTE takes inspiration from both Task2Vec [5] and OTDD [11], it creates a dataset embedding (Task2Vec inspiration) based on solving an OT problem (OTDD inspiration). Like OTDD, WTE is model-agnostic and free of training, capable of handling (partially) disjoint label sets. Like other embedding-based dataset similarity measures, WTE alleviates the problem of expensive pairwise computation by employing embeddings as task representations. The WTE associated similarity correlates highly with the OTDD distance while reducing the computational cost from quadratic to linear with respect to the number of tasks. Like OTDD, WTE represents labels as their feature-based distributions, using Gaussian approximations to speed up computation. Additionally, WTE embeds these label distributions through multi-dimensional scaling (MDS), a non-linear dimensionality reduction approach preserving pairwise distance. Afterward, the MDS-based label embeddings are concatenated to the feature vectors, yielding the updated samples. Tasks' updated samples are embedded using a, previously proposed, Wasserstein embedding method. The Wasserstein embedding creates an embedding for probability distributions, for which the Euclidean distance between them approximates the 2-Wasserstein distance. Hence, the distance between tasks is the 2-Wasserstein distance between their updated samples, which is approximated by the (squared) Euclidean distance between the embeddings.

2.2.1.3 Configuration Characterization Methods

RankML’s Configuration Characterization The authors of [68] introduce RankML, a meta-learner employing dataset and configuration characterizations using a novel ML pipeline representation approach. This configuration characterization method represents the topology of the ML pipeline as a sequence of words. Following our formalization (see Definition 1), each $a \in \mathbf{A}$ and its accompanying hyperparameters in λ are represented by a unique fixed-length hash, with a sequence thereof composing the configuration characterization. See Figure 2.3 for example configuration characterizations representing their respective pipeline. Note that the method employs the hyperparameters λ in pipeline (g, \mathbf{A}, λ) to characterize the pipeline while simultaneously yielding a compact pipeline representation. Adversely, the unique hashes composing the compact pipeline representation together with a huge search space imply many different elements in the pipeline representation, which may be challenging for a meta-learner to distinguish. The pipeline configuration is created using the following rules:

1. Define the configuration characterization’s dimensionality as the longest pipeline to characterize.
2. Hash each node (algorithm a and its accompanying hyperparameters in λ) in the pipeline in reverse order (from output to input).
3. Each node in g must be hashed before its inputs.
4. If there are parallel sub-pipelines, the longest sub-pipeline is processed first, ties are broken randomly.
5. All pipeline characterizations have equal dimensions, pad with hashes representing "blanks" if necessary.

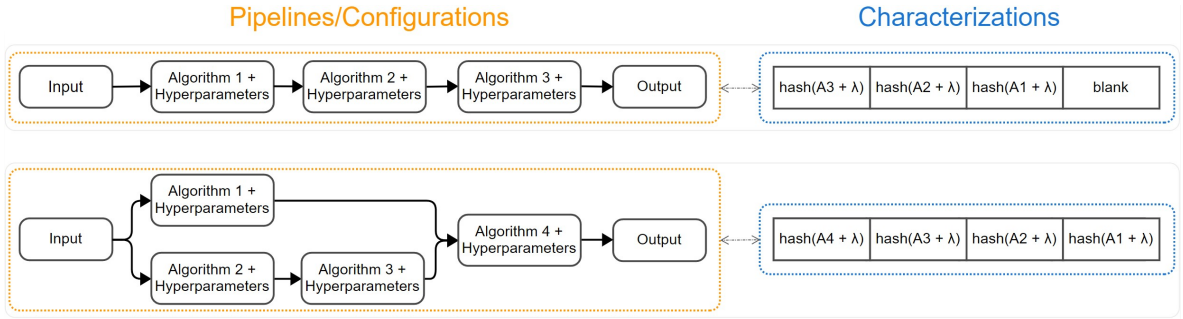


Figure 2.3: Example depicting configurations (i.e. pipeline structure g with its algorithms A and accompanying hyperparameters λ) and RankML’s corresponding characterizations following aforementioned rules (1, 2, 3, 4, 5). The example is modified from [68].

Decomposition-Based Configuration Characterizations Probabilistic Matrix Factorization [45] (PMF) is a collaborative-filtering-inspired AutoML approach considering a matrix $\mathbf{Y} \in \mathbb{R}^{N \times D}$ composed of dataset-pipeline evaluations for N pipelines and D datasets. PMF seeks a low-rank decomposition $\mathbf{Y} \approx \mathbf{X}\mathbf{W}$ with $\mathbf{X} \in \mathbb{R}^{N \times Q}$ and $\mathbf{W} \in \mathbb{R}^{Q \times D}$ with Q denoting the dimensionality of the learned latent space. The goal of PMF is to automate the selection and tuning of pipelines consisting of a pre-processing step and a predictive model, but in creating the low-rank decomposition it additionally learns to embed pipelines in the latent space purely from their performance in \mathbf{Y} . The learned matrix \mathbf{X} thus contains the N configuration characterizations as Q -dimensional vectors in the learned latent space. The approach is computationally demanding, because it requires decomposing a high-dimensional matrix with PCA, where the computational cost grows cubically with the number of encountered datasets and quadratically with pipelines (whose configuration space is infinite).

Like PMF, TensorOBOE [138] uses a low rank decomposition for pipeline search, but on tensors instead of matrices. The 6-order error tensor ϵ contains prior pipeline-dataset evaluations, where each direction represents a different aspect, namely: datasets, imputers, encoders, standardizers, dimensionality reducers and estimators. We refrain from repeating elaborate technical details in this high-level overview, for more details consider Section 3.3 of the paper. In short, the tensor is decomposed using Tucker decomposition to yield one core tensor and six factor matrices (one per direction in ϵ). The columns (1 for each matrix) together embed a pipeline, which composes the configuration characterization. Clearly, computing the Tucker decomposition of a 6-order tensor is even more computationally expensive than computing a PCA-based matrix factorization, for details on the time complexity of low-rank Tucker decompositions consider [82]. Though the time complexity may differ per solver it is safe to say that the required computational resources are beyond that of AutoML systems, with TensorOBOE requiring a 128 core machine with 1056GB of memory for 215 small-sized datasets and 23.424 pipelines.

2.2.2 Meta-Learners

A meta-learner is a component performing knowledge transfer from prior to new tasks in a meta-learning system. In performing knowledge transfer a meta-learner may require information regarding the learning task or prior solutions. Therefore, we can consider three groups of AutoML-related meta-learners, namely; similarity-based meta-learners which employ

dataset similarity; characterization-based meta-learners which employ dataset or configuration characterizations; and dataset-agnostic meta-learners which do not depend on datasets. Thus, a meta-learner could simultaneously be similarity- and characterization-based. In the following subsections we give an overview of the state-of-art meta-learners categorized according to our previously defined groups. Meta-learners employing multiple components simultaneously are discussed in the Section whose component’s usage is most pronounced.

Scoping Due to the broadness of the meta-learning research field, a meta-learner can perform a wide variety of tasks including those in few-shot learning, reinforcement learning and deep learning [52]. The scope of our meta-learner discussion concerns approaches specifically operating on tabular datasets (see Restriction 2) such that these can be used for tabular AutoML systems. The aforementioned application fields tend to be neural-network driven and therefore tailored to data modalities different from tabular data, causing their meta-learning approaches to be excluded from the current literature review.

In the overview we focus on works that most closely follow our formulation and formalization of the AutoML problem (see Section 2.1), considering the *pipeline* creation problem. Meta-learners solely considering a subproblem of the pipeline creation problem, i.e. only selecting the pipeline structure, or its algorithms or hyperparameters are out-of-scope because their incorporation in complete-pipeline AutoML systems is less straightforward. Likewise, meta-learning approaches crafted solely for data preparation [143, 17, 49, 65, 18] or feature generation [59] are excluded from this overview. For similar reasons, we disregard meta-learning and AutoML approaches opting for multiple stages, such as a bi-level scheme [145], two-stage approach [105] or even more stages as in [87]. The latter distinction dismisses meta-learning approaches within multi-stage settings, including [148, 88, 107].

Lastly, we exclude approaches that explicitly rely on new or specific prior dataset-pipeline evaluations, because it limits their incorporation in AutoML systems. Despite a fine line, we argue works (iteratively) gathering new evaluations are better classified as AutoML systems instead of meta-learning approaches. Compared to meta-learning approaches, entire AutoML systems are limited with respect to their modularity, in turn restricting their AutoML adoption and causing their exclusion from this overview. For similar reasons, works that assume a specific structure for their metadataset are excluded as well.

We thus exclude approaches developed with a specific evaluation scheme in mind, that being prior to pipeline synthesis or during. Exclusions include approaches performing a matrix decomposition such as Probabilistic Matrix Factorization (PMF) [45] and TensorOBOE [138] because they require pipeline evaluations for a novel dataset and gather new evaluations thereafter. Additionally, PMF explicitly assumes the metadataset is composed of a set of previously defined possible pipelines. Other exclusions are the meta-reinforcement-learning-based AlphaD3M [32] and a phased performance-based pipeline planner P4ML [48], because both require pipeline evaluations in their setup.

To summarize we exclude any meta-learning approach: not operating on tabular data, specifically crafted for (pipeline) sub-problems, considering multi-phase setups, gathering new evaluations or imposing metadataset assumptions. Our scope is supported by a desire for a modular meta-learning framework (see Section 1.2), AutoML research increasingly focusing on complete ML pipelines (for tabular data) [147], and pre-existing literature already comparing meta-learning approaches for subproblems [60].

2.2.2.1 Similarity-Based Meta-Learners

Similarity-based direct configuration transfer Configuration transfer applications reuse configurations from (similar) previous tasks for the current task. In reusing previous configurations an inductive bias is transferred, reducing the effort to choose among a vast amount of possible configurations. The simplest similarity-based configuration transfer approach is to use the best-performing configuration from the most similar task. Though this likely yields a non-optimal configuration, its application in an AutoML tool for time-series data [66] shows promising configurations can be identified. Another option is to evaluate several recommended configurations [126], transferring the best evaluated configuration.

Similarity-based direct configuration transfer for warm-starting optimization Besides directly using other tasks’ configurations, there are also opportunities to use them indirectly by warm-starting optimization for the current task. The goal of warm-starting is to increase convergence rate and/or improve final performance by selecting configurations from previous tasks to yield a high-quality initialization for the optimization procedure. In this setup, the role of a dataset similarity measure is to identify the prior tasks whose configurations are transferred. In doing so, it is assumed that similar tasks have similar high-performing configurations.

One approach, as presented in [43], is to use the t best configurations from the d most similar tasks, yielding $t \cdot d$ configurations for initialization. Because we need to evaluate the $t \cdot d$ initial configurations, we need to be mindful of this value; it directly influences the exploration-exploitation trade-off of AutoML systems. If $t \cdot d$ is too small, we may under-exploit and could have found better configurations through a better initialization. If $t \cdot d$ is too large, we waste the exploration potential of the optimization procedure. In the extreme case, with $t \cdot d$ being the total amount of evaluations, we again yield a direct configuration transfer mentioned previously. Not only $t \cdot d$ influences this trade-off, but also the individual choices for t and d ; a higher t resembles more exploitation whereas a higher d promotes exploration.

To the best of our knowledge, there is no consensus on choosing t and d , nor are there other approaches to incorporate similarity in selecting configurations for warm-starting pipeline optimization. There is however much related work on warm-starting hyperparameter optimization (HPO) using similarity information with approaches for Bayesian Optimization [42, 125, 61] and CMA-ES [96] showing performance increases. Since pipeline optimization is more complex than HPO, warm-starting pipeline optimization is especially promising with [43] noticing complex problems benefiting most from warm-starting Bayesian optimization.

A critical reader may have noted that the discussed approach [43] is applied on HPO, while we excluded meta-learners for sub-problems from the discussion’s scope. The meta-learning approach is included because it has been successfully applied in warm-starting search for entire pipelines in subsequent work [41]. For that reason, similar meta-learning approaches (e.g. [134]) without such follow-up works are not discussed.

Similarity-based search space pruning for warm-starting optimization Apart from direct configuration transfer, there are other approaches for warm-starting optimization procedures based on dataset similarity. We can transfer knowledge not only by promoting certain parts of the search space, but also by excluding parts thereof. This core idea – pruning the search space for an AutoML system’s optimization procedure based upon a similarity measure – is investigated in [94] for the AutoWeka4MCPS [114] AutoML system. The core idea is not

novel, for it had already been explored in [133], but the extension to a more complete setting is, i.e. warm-starting full pipeline search instead of HPO.

Following our notation (see Definition 1), [94] prunes the search space (initially $|\mathcal{A}| = 30$) through excluding all but the k best algorithms $a \in \mathcal{A}$, thus yielding $|\mathcal{A}| = k$. Interestingly, the paper compares this meta-learning approach in a setting employing dataset similarity – only considering the performance on the most similar datasets – and in a dataset-agnostic setting, but can only affirm the former approach to be significantly better than the baseline (and is therefore treated here instead of Section 2.2.2.3).

It is worth noting the dataset-similarity measure is modular with respect to the meta-learning approach. The authors opt for a meta-feature-based dataset-similarity measure solely employing landmarking meta-features, begging the question whether these results can be improved with more diverse meta-features or learned dataset characterizations.

The experimental evaluation explores the effect of choosing k , which we can consider a exploration-exploitation trade-off parameter, with decreasing k exploiting more and increasing k exploring more. The findings indicate that among $k \in \{1, 4, 8, 10, 19\}$ setting $k = 10$ is optimal using the similarity-based approach. Interestingly, the choice for k and thus the exploration-exploitation trade-off, differs for the dataset-agnostic and dataset-similarity-based meta-learning approaches, with the higher-performing approach preferring a higher value for k . This discombobulating finding suggests increased meta-learner performance need not imply favoring exploitation over exploration in warm-starting AutoML search. Further complicating the trade-off choice, the value for k depends on the subsequent optimization procedure’s time budget, with a higher time budget allowing for more exploration.

2.2.2.2 Characterization-Based Meta-Learners

RankML RankML [68] ranks machine learning pipelines using a XGBoost-based ranker trained on dataset- and configuration characterizations together with their associated performance. The pipeline configuration is characterized through a novel representation of its topology (see Section 2.2.1.3) while descriptive and correlation-based meta-features characterize the data. RankML employs distinct XGBoost ranker models per task (regression or classification) and evaluation metric. By default RankML ranks all pipelines in the metadataset on the novel task to recommend the top ranked pipeline without the need for evaluation. However, better performance is achieved by selecting, through evaluation, the best-performing pipeline from a set of top-ranked pipelines. If RankML recommends the best-evaluated top-4-ranked pipeline, its performance is on par with TPOT [98] and Auto-Sklearn [40], but at reduced computational cost.

Similar to many meta-learners, a disadvantage of RankML is its poor ability to extrapolate beyond the metadataset; RankML solely ranks previously-created pipelines, it does not originate them. The authors acknowledge this weakness, but argue a sufficiently large metadataset (i.e. their 313.488 pipeline evaluations) partially mitigates the issue. Note the RankML meta-learner need not be tailored to specific dataset nor configuration characterization methods. Correspondingly, according to the authors, investigating whether the RankML setup benefits from more recent dataset characterizations (see Section 2.2.1.2) is an interesting research direction.

The use of RankML in AutoML systems is investigated in MetaTPOT [69], specifically exploring the incorporation of a ML pipeline ranking model in the synchronous evolutionary AutoML system TPOT. The work shows MetaTPOT attains decreased TPOT runtime without

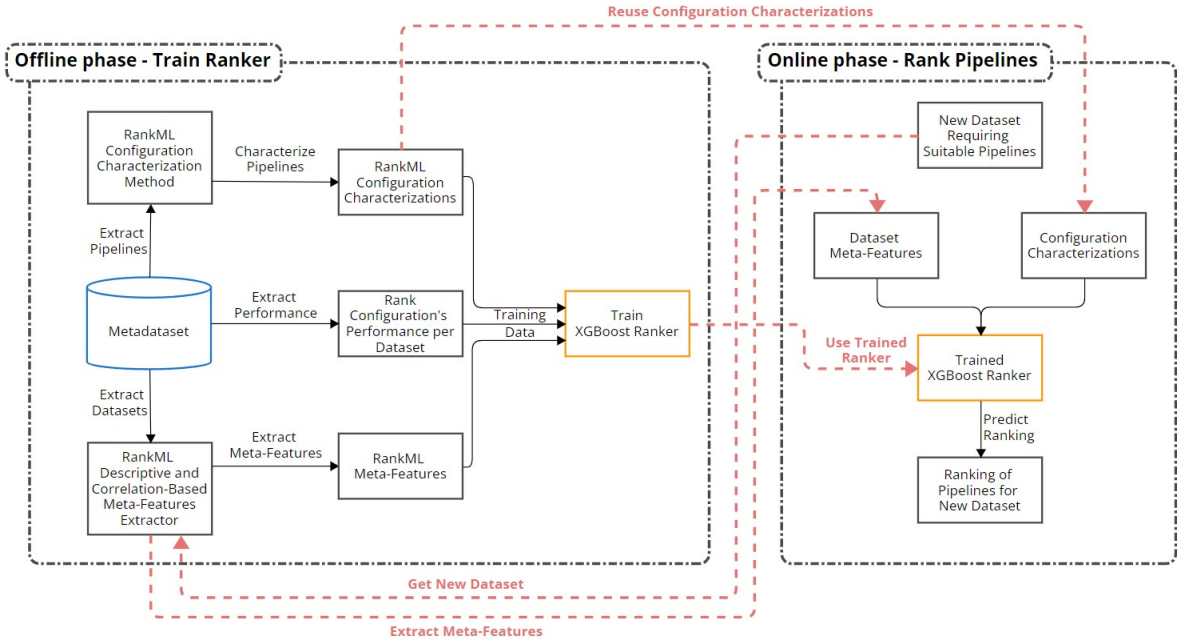


Figure 2.4: Illustration of the RankML [68] meta-learner, arrows illustrate the steps and red arrows highlight interactions between the offline and online phase. The left part of the figure depicts the meta-learner offline phase in which the XGBoost-based ranker is trained using dataset and configuration characterizations, with their associated ranks. Datasets are characterized using descriptive and correlation-based meta-features, pipelines are characterized using the RankML configuration characterization method (see Figure 2.3), and their performance is ranked per dataset. The right part of the figure displays the online phase, in which the metadataset pipelines (candidate pipelines) are ranked for a new dataset using the trained ranker from the offline phase. The input for the ranker consists of a characterization of the new dataset and the configuration characterizations previously computed in the offline phase.

negatively affecting performance. MetaTPOT does not exhibit improved accuracy-related performance, which we postulate may be attributed to a limited and non-uniform incorporation of pre-ranked individuals in warm-starting the evolutionary search procedure.

2.2.2.3 Dataset-Agnostic Meta-Learners

Portfolio building Portfolio building is a dataset-agnostic meta-learning approach introduced in [36] and incorporated in auto-sklearn 2.0 [38] to replace the original meta-features based approach. The approach uses prior evaluations to create a portfolio \mathcal{P} , an ordered set of pipelines (g, \mathbf{A}, λ) , of a specified size $|\mathcal{P}|$ that are expected to perform well, regardless of the new dataset. The approach is intended to warm-start a subsequent optimization procedure, but with $|\mathcal{P}|$ as a parameter this need not be the case (e.g. $|\mathcal{P}| = 1$).

In an offline setting, the approach commences by searching the best configuration for each dataset $d \in \mathcal{D}$ to create the set of candidate pipelines \mathcal{C} for the to-be-built portfolio. Each configuration $c \in \mathcal{C}$ is evaluated on each dataset $d \in \mathcal{D}$ using a cross-validation procedure to yield performance matrix \mathbf{P} , a $|\mathcal{D}| \times |\mathcal{D}| = |\mathcal{C}| \times |\mathcal{D}|$ dimensional matrix. Performance matrix \mathbf{P} is transformed to error matrix \mathbf{E} with equal dimensionality by computing the normalized regret across all datasets.

Portfolio \mathcal{P} is built by employing greedy submodular function minimization [64] on \mathbf{E} , with a process as follows. The portfolio is built iteratively by greedily selecting configurations one by one: in each iteration, we add the candidate $c \in C$ to portfolio \mathcal{P} such that the portfolio's generalization error is minimized. The generalization error is the sum of the regret on each dataset for its highest-performing configuration in $|\mathcal{P}|$. If multiple candidates reduce the generalization error equally – with a special, yet not uncommon, case being no reduction – then the first evaluated candidate is added to \mathcal{P} . The portfolio building procedure stops when the portfolio size is reached, which is upper-bounded by $|\mathcal{D}|$.

In a warm-starting scenario, portfolio size $|\mathcal{P}|$ acts as an exploration-exploitation parameter, with increasing values exploiting more, and decreasing values exploring more. If we were to generalize beyond the introduced framework – considering C to not only consist of the best configuration per dataset d , but the best k configurations – then k would act as a trade-off parameter similar to $|\mathcal{P}|$. Moreover, in this generalized setting the upper bound on the portfolio size increases by factor k .

Chapter 3

MLTA: a Meta-Learning Toolbox for AutoML

In this chapter we detail the proposed Meta-Learning Toolbox for AutoML (MLTA). In Section 3.1 we employ the preliminary knowledge from Section 2.2 – especially the overviews on dataset characterization methods (Section 2.2.1.2), dataset similarity measures (Section 2.2.1.1), configuration characterization methods (Section 2.2.1.3) and meta-learners (Section 2.2.2) – to select meta-learning components to incorporate in MLTA and compare in the experimental evaluation (see Chapter 4). In Section 3.2 we present the (modular) structure and functionality of the toolbox, detailing which meta-learning approaches can be built using its main components: dataset characterization methods, dataset similarity measures, configuration characterization methods and meta-learners. In addition to that, we perform experiments analyzing the run time for dataset characterization methods in Section 3.2.2.

3.1 Meta-Learning Component Selection

In this section we compare and select approaches from the preliminary Chapter (2) to incorporate in MLTA, sometimes generalizing beyond their initial interpretation. We explicitly defer from (re-)explaining the approaches, because they are already explained in Chapter 2. The approaches to compare are few in number and relatively well-fitting due to extensive scoping in the preliminary Chapter, guided by Restrictions 1, 2, 3, 4, 5 and the aim for a modular framework. Nonetheless, we need to select – per meta-learning component – which approaches are to be incorporated in the toolbox. This selection is guided by a comparison on a set of key dimensions, namely: applicability, scalability and utility. The materialization of these properties for the different meta-learning components (dataset characterization methods, dataset similarity measures, configuration characterization methods and meta-learners) will be discussed in their respective subsections (see Sections 3.1.1, 3.1.2, 3.1.3, 3.1.4). To permit comparing approaches, we will argue for the most relevant characteristics of each dimension under consideration, which we term aspects of each dimension.

3.1.1 Dataset Characterization Method Selection

In this subsection we select dataset characterization methods for MLTA guided by a comparison on applicability, scalability and utility of the approaches. It is important to note that, likely,

there is no one-size-fits-all dataset-characterization method, because what constitutes the most suitable method may be dependent on the problem at hand. One must consider the measures' applicability and there is a trade-off between computational cost and utility, which are both dependent on the data and task to be solved. In the following paragraphs we will argue for the aspects of each dimension, they are as follows. For the dimension of applicability we argue for the aspects of data modality and task dependency. The scalability dimension's aspect concerns scalability with respect to the data. The last dimension, utility, has the main aspects of robustness, parameterization and effectiveness.

Applicability The applicability dimension concerns requirements for the ability to compute a dataset characterization. The applicability of a dataset characterization method depends on its input, namely the data representing a task to be solved. Therefore, we evaluate the applicability with respect to data and tasks. A dataset may vary widely, on lots of different facets, but possibly, the most informative and limiting aspect of a dataset is its modality. We can distinguish between many data modalities, but consider only the most common: image data, tabular data, and text data. We aim for a method that is at least applicable on tabular data, without restrictions on its schema. The second aspect of applicability concerns the task dependency which – due to having restricted the analysis to a supervised setting – are binary or multi-class classification or regression tasks. We aim for methods supporting both binary and multi-class classification tasks, where the additional applicability on regression tasks would be welcome.

Scalability Scalability concerns the ability to perform the method efficiently and in a tractable way, involving base costs and costs scaling with dataset size. We deem a dataset characterization method scalable, with respect to the data, if its computational time complexity is at most linear with respect to the number of samples, features, and outcomes.

Utility The utility dimension is concerned with the use of a dataset characterization method or its output. Therefore, we can distinguish between aspects concerning the production of the output, or the output itself. Given equal input, in the form of datasets, there are two aspects influencing the production of the output of a task-similarity measure: the measures' robustness and its parameterization. Additionally, the quality of the dataset characterization is important. However, with a lack of ground truth we can only assess it in a behavioral way: the effectiveness, which is highly dependent on the data and task at hand, especially since there are many different downstream applications. Due to the need for a behavioral/experimental evaluation of effectiveness we exclude it from the comparison.

Robustness is the stability of the output, given the same – or perhaps slightly different – input. Hence, a robust method achieves stable characterizations invariant to meaningless changes in the input. A dataset characterization method takes a dataset as input, sets are *unordered* collections of items. Consequently, the permutation of the samples of a dataset is a meaningless change in the input. Hence, for a dataset characterization method to be robust it must be invariant to the permutation of its samples. Following a similar line of reasoning, when features are independent of each other, then the measure should be invariant with respect to the permutation of features, but only if the permutation is consistent across all samples. Robustness also involves stability of the output when there is no change in the input. A non-deterministic method is not necessarily undesirable, as long as its output is stable. Thus

the degree of robustness of a dataset characterization method involves: invariance to sample order; invariance to feature order (if the feature order is independent of the outcome), and a stable output given equal input.

Types of parameterization come in two forms, namely methods that are parametric or non-parametric. Parametric methods are more difficult to use, because the output and its effectiveness depend on user input. Therefore, especially in the context of automation, non-parametric dataset characterization methods are preferred.

In the remainder of this Subsection, we assess the dataset characterization methods presented in Section 2.2.1.2 on the aforementioned dimensions and their aspects, for a brief overview consider Table 3.1. Subsequently, we use these aspects to compare and select approaches for MLTA.

Meta-Feature-Based Dataset-Characterization Method The properties of meta-feature-based dataset characterizations and accompanying similarity measures depend on the collection of meta-features used. They can be applicable to multiple data modalities (including text, image and tabular data) and multiple tasks such as regression, classification and clustering. Moreover, they can be scalable and fast to compute (mostly simple meta-features) and robust without the need for parameterization. Contrarily, they can also be tailored to a data modality and task type, computationally demanding, require parameterization and lack robustness.

For comparison sake, we determine the properties of the common meta-feature collections by Feurer et al. [43] and Wistuba et al. [134] previously discussed in Section 2.2.1.2 and detailed in Appendix C. These meta-features are applicable to image and tabular datasets in binary and multi-class classification settings.

Both collections tend to be computationally undemanding, but this depends on the dataset at hand. Being composed of simple and low-complexity statistical meta-features, the Wistuba collection is scalable with respect to number of instances, features and outcomes and therefore the data. Contrarily, the Feurer meta-feature collection requires the more computationally demanding linear discriminant analysis (LDA) and principal component analysis (PCA), whose time-complexity can grow poorly depending on the solver, but is always non-linear for either features, or instances or outcomes. Moreover the Feurer meta-features include landmarking models such as KNN, growing quadratically in the number of samples. Thus, due to the incorporation of PCA-based, LDA-based and landmarking-based meta-features the Feurer meta-feature collection is not scalable with respect to the data, potentially leading to a poor run time on larger datasets.

Regarding utility, both meta-feature and summarization function collections do not require further parameterization. The robustness for the Wistuba meta-features is excellent due to a deterministic computation of the meta-features, invariant with respect to meaningless permutations in the instances, features and outcomes. Contrarily, the Feurer meta-features are less robust due to its additional PCA-based, LDA-based and landmarking-based meta-features. Both the tree-based landmarking models and LDA are non-deterministic methods, with the latter *possibly* being sensitive to the data at hand [62]. Therefore it is debatable whether the Feurer meta-features produce a stable dataset characterization. However, many of these features are invariant with respect to meaningless permutations in instances, features and outcomes. Therefore, we opt for classifying the Feurer-meta-features-based dataset characterization to be semi-robust; seemingly there are no prevalent robustness issues, though an experimental evaluation is required to confirm this.

Task2Vec [5] Task2Vec’s use of a probe network determines some of its properties. First of all, the probe network is considered a parameter, with the paper itself showing considerable performance differences based on its choice. The CNN-based probe network(s) also determine the data modality to be images, with a requirement for their size. Hence, the approach may be less applicable to images that cannot be properly scaled to the pre-defined resolution. Moreover, the probe networks considered are specifically pre-trained on a classification task. The authors argue the method also computes meaningful embeddings for regression if the probe network performs a regression task, but the experimental evaluation thereof is lacking.

Task2Vec is not highly robust for several reasons involving the probe network. Computing the embedding involves re-training the probe network’s classification layer. Therefore, the order of the instances influences the computation of the embedding. Moreover, neural network training and the estimation of the FIM are non-deterministic, possibly resulting in an unstable characterization. The computation of the embedding involves both re-training on the task and FIM estimation. The estimation of the FIM is independent of the number of samples, their resolution, and the outcomes. Moreover, the time complexity for training a neural network grows linearly in the number of samples and constant in the number of outcomes. Therefore, although the base cost may be high, Task2Vec scales well with increasing data size.

Dataset2Vec [58] Dataset2Vec is most suited for tabular datasets because it learns the relation between single predictor-target pairs. Due to this setup, Dataset2Vec likely yields poor performance on other data modalities. For instance, for image data Dataset2Vec would consider the relation between specific pixels and the target. It would completely disregard the spatial structure necessary for meaning. Due to the setup of the pre-trained neural network Dataset2Vec is only suited for classification datasets. To the best of our knowledge, there currently is no effort extending Dataset2Vec to a regression problem setting.

Because Dataset2Vec only needs to pass a specified number of batches through the pre-trained network to compute the characterization it is scalable with respect to the data. Moreover, Dataset2Vec is feature, sample and outcome invariant because it uses the coordinates in the pre-trained network’s latent space directly, which only depends on the batches. However, the batching procedure causes the method to be non-deterministic, possibly yielding unstable characterizations. Luckily, this issue is mitigated by setting a seed for the batching procedure. Therefore, we can still claim that DataSet2Vec is a robust method. However, the characterization – and especially its accuracy and validity – depends on the number of batches that are passed through the pre-trained network. Though the size of the batches is determined by the pre-trained network, the number of batches is a parameter (with a default value of 10 in the original work). Unlike Task2Vec, the neural network employed by Dataset2Vec is not a parameter, because Dataset2Vec directly uses coordinates in the learned pre-trained neural network’s latent space as a dataset representation.

DIDA [26] As shown in its paper’s experimental evaluation, architectural variants using DIDA components can be applied to both regression and classification tasks. In essence, DIDA can be applied to any dataset with features and outcomes. However, because it does not take into account spatial structure, it is best applied to tabular datasets. DIDA does not rely on a pre-trained model to extract meta-features, they need to be learned and thus computed each time. The computational time complexity scales quadratically in the number of samples, and therefore DIDA is not scalable with respect to the data. Though DIDA follows a similar

setup as Dataset2Vec – characterizing a dataset using coordinates in a trained network’s latent space – it is not scalable because it does not employ a pre-trained model during dataset characterization inference, it needs to be trained.

The main selling points of the approach are its invariance properties with respect to the data. Moreover, confidence estimates in the experimental evaluation suggest that the output is sufficiently stable. Concluding from the invariance properties and the characterization stability in their experimental evaluations we consider DIDA to be robust. Lastly, DIDA is parametric because learning meta-features for one dataset requires training on a patch identification task, involving another dataset to be specified.

WTE [77] Due to a setup similar to OTDD, the WTE dataset characterization can be computed for tabular data, image data and embedded text data. Similarly, due to explicitly modeling labels the method’s applicability is restricted to classification tasks. The computational time complexity of the WTE characterization method grows cubically with respect to the number of samples, it is thus not scalable with respect to the data. Unlike OTDD – inheriting its invariance properties from the optimal-transport setup – WTE is not invariant to permutations of its samples. Recall WTE computes label embeddings by multidimensional scaling (MDS), MDS is not invariant to sample permutation and therefore neither is WTE. Whether the characterization is stable is debatable, we only know the associated similarity value correlates highly with OTDD, which itself is stable as shown in their experimental evaluation. Thus to claim stability of the characterizations one must assume that a stable similarity value implies a stable characterization and that a highly correlated similarity value implies stability thereof. We deem both these assumptions strong and not at all obvious and therefore, together with not satisfying invariance, we classify the WTE approach to not be robust. Lastly, we note that the dimensionality of the MDS output – which encodes the label distribution – is a parameter with a default value of 10.

Comparison and Selection The comparison and selection are guided by the approaches’ aspects depicted in Table 3.1, therewith we can easily distinguish between approaches. First of all we consider the applicability dimension; we can compare the methods on their applicable data modality: suggesting the preclusion of Task2Vec, because it only operates on image data. Moreover, its exclusion is further supported by its poor utility. Secondly, we cannot discard any characterization method based on their task dependency, because all methods support both binary and multi-class classification.

The following dimension under consideration is scalability. We can distinguish approaches based on whether they need to train/learn a model *on the spot*, either for learning a dataset characterization or not. Approaches learning dataset characterizations are to descriptive meta-features as neural networks are to classical features, that is: they learn a way to represent data instead of hand-crafting a representation. Like training neural networks, learning dataset characterizations seems to be a computationally intensive task; none of these approaches are scalable. Note that Dataset2Vec is scalable even though it has learned a dataset characterization. It achieves this because it employs a pre-trained network and therefore need not learn the dataset characterization on the spot. In similar fashion, by using a pre-trained network, Task2Vec also achieves scalability with respect to the data. Perhaps surprisingly, we do not exclude approaches based on their poor scalability with respect to the data. In contrast to a poor utility or applicability, a poor scalability need not imply an infeasible method, though it

does limit the scenario's for which it is feasible. In other words, an unscalable method may still be used for smaller datasets. Therefore, we do not exclude Feurer meta-features, DIDA and WTE purely on this aspect.

Next, we consider the utility dimension for the remaining approaches: Wistuba Meta-Features (WistubaMF), Feurer Meta-Features (FeurerMF), Dataset2Vec, DIDA and WTE. Observe we can distinguish the approaches on parameterization by whether or not they learn a dataset characterization. We exclude WTE based on its poor robustness in combination with its poor scalability and the need for parameterization. We exclude DIDA due to its daunting parameterization: a difficult-to-choose supplementary dataset for which – in the original work – there is no sensible default nor a proper argument to select it. The remaining approaches attain sufficient utility, because they do not have poor robustness (FeurerMF is not clear), nor difficult-to-choose parameters. We do not exclude Dataset2Vec based on the parameterization aspect, because the original work provides a sensible and well-performing default value which we need not alter.

Lastly, we consider supplementary arguments for excluding the approaches to further strengthen our selection. In that regard, WTE is not well suited for characterization-based meta-learners because it does not produce fixed-size characterizations.

By comparing the dataset characterization approaches on their main dimensions and their corresponding aspects we have selected the following methods for incorporation in MLTA: WistubaMF, FeurerMF and Dataset2Vec. We note that both WistubaMF and FeurerMF are strongly related methods, and could be considered different instantiations of a meta-feature-based characterization method. Nonetheless, we include both approaches because they satisfy our inclusion criteria. Moreover, since the meta-features in WistubaMF are a subset of those in FeurerMF we are eager to compare them in an experimental setting (see Chapter 4).

3.1.2 Dataset Similarity Measure Selection

In this subsection we select dataset similarity measures for MLTA guided by a comparison on applicability, scalability and utility of the approaches. It is important to note that, akin dataset characterization methods, there is no one-size-fits-all measure, because what constitutes the most suitable method may be dependent on the problem at hand. The materialization of the dimensions for a dataset similarity measure is akin to that of a dataset characterization method, they are as follows.

For the dimension of applicability we argue for the aspects of data modality and task dependency. The scalability dimension concerns scalability with respect to the data *and the number of datasets*. The last dimension, utility, has the main aspects of robustness, parameterization and effectiveness. Note that these aspects are akin to the aspects for a dataset characterization method, which should not surprise with the existence of characterization-based similarity measures. The main difference is on the scalability dimension; for dataset similarity measures it is important that they are scalable with respect to the number of datasets to compute the similarity value for. In addition to that the remaining aspects are different because they now need to be applied to different input – two datasets instead of one – and a different output – a similarity value instead of a characterization. Therefore, we need not explain why the remaining aspects are suitable for a dataset similarity measure, we only need to explain what constitutes the scalability dimension.

Scalability concerns the ability to perform the method efficiently and in a tractable way, involving both the base cost of performing the similarity measure once, and scalability to

	Applicability		Scalability		Utility	
Char. Method	Data modality	Task dependency	Data	Robustness	Parametrization	
WistubaMF [134]	Tabular, Image	Classification	✓	✓	✓	None
FeurerMF [43]	Tabular, Image	Classification	✗	~	✓	None
Dataset2Vec [58]	Tabular	Classification	✓	✓		Number of batches
Task2Vec [5]	✗ Image	Classification, Regression	✓	✗		probe network, α : complexity
DIDA [26]	Tabular	Classification, Regression	✗	✓		Supplementary dataset
WTE [77]	Image, Tabular, Embedded Text	Classification	✗	✗		MDS dimensionality
Sim. Measure						
OTDD [11]	✗ Image, Embedded Text	Classification	✗	✓		Feature distance metric
WTE [77]	✗ Image, Embedded Text	Classification	✗	✗		MDS dimensionality

Table 3.1: Comparison of dataset characterization methods and dataset similarity measures on applicability, scalability, and utility. Note that each fixed-dimensional dataset characterization method (all but WTE) has an associated characterization-based dataset similarity measure with the displayed properties. WTE is presented twice because its aspects differ for its different uses. A green check mark for an aspect of the dimension indicates the corresponding approach attains its properties in desirable fashion, a red cross implies it does not, and an orange tilde represents semi-attainability.

multiple tasks, thus performing it multiple times. We deem a dataset similarity measure scalable with respect to the data, if its computational complexity is constant with respect to the number of samples, features, and outcomes for *both datasets*. Moreover, similarity measures should be scalable with respect to the number of datasets (e.g. tasks), because it is important for similarity-based meta-learners to compare the similarity among many datasets. We deem a dataset similarity measure scalable with respect to tasks, if the amount of computational resources involved in computing all pairwise similarities between a group of tasks grows approximately linearly.

An ideal dataset similarity measure should be applicable to two, possibly different, datasets, each allowing for any combination of data modality and task type. In addition to that it should be scalable with respect to the data and number of datasets. Lastly, the computation of the similarity value should be robust and without requiring the specification of difficult-to-select parameters.

In the remainder of this Subsection we assess the dataset similarity measures presented in Section 2.2.1.1 on the aforementioned dimensions and their aspects. We additionally include all possible characterization-based dataset similarity measures (see Section 2.2.1). Recall, these

characterization-based similarity measures simply compute the (dis)similarity directly from the characterization and are therefore only possible if their corresponding characterization method produces fixed-size characterizations. This consideration excludes WTE to have a characterization-based similarity measure because its characterization dimensionality is dataset dependent. All characterization-based dataset similarity measures are scalable with respect to the number of datasets because for N datasets we only need to compute N characterizations to compute N^2 pair-wise similarity values. All other aspects are directly inherited from the corresponding characterization method (see the top part of Table 3.1).

OTDD [11] Several of OTDD’s aspects follow from it assuming access to a distance metric for feature vectors. Firstly, it makes the distance metric for feature vectors a parameter. Secondly, because of abstracting the distance metric away, it makes the method feature invariant. Thirdly, it disallows the direct application of OTDD on any dataset without a proper distance metric for feature vectors. However, distance metrics for vectors of equal size are well-known and therefore OTDD can be applied to datasets with feature vectors of equal dimensions. This restriction does prohibit most practical applications to tabular datasets, because their feature vectors typically vary in dimensionality.

Considering the suitable data modality, OTDD is applicable to text-based and image-based data, but the text needs to be encoded and the images need to be rescaled to equal resolution. Hence, we deem OTDD to be applicable with respect to images and embedded text because their equal dimensionality between datasets is not a strong assumption. Contrarily, assuming equally dimensional features for tabular datasets is a strong assumption, violating desired schema agnosticism, which leads us to exclude it from OTDD’s applicable data modalities. Regarding task dependency, OTDD explicitly models label distributions, therefore it is only applicable to classification tasks.

Regarding scalability, OTDD with the computational speed-up still scales cubically in the number of features and quadratically in the number of samples. Therefore, it is far from scalable with respect to the data and thus best used on datasets with a small number of samples and features. Moreover, it implies that images must be of low resolution and text embeddings must have low dimensionality. OTDD computes a pair-wise distance between two datasets without an intermediary task representation, such as an embedding or characterization. Therefore, OTDD must be fully recomputed for each pair of datasets, disallowing it to scale well with respect to the number of datasets.

Regarding utility, the authors show that the stability of OTDD’s output increases with an increasing number of samples, with it also being stable overall. Furthermore, due to the optimal transport setup, the method is sample and feature invariant. Because OTDD attains invariance properties and because its output is stable it can be considered robust.

WTE [77] Due to a similar setup as OTDD, WTE is only applicable for image data and embedded text because it requires equally-sized features across datasets. It thus suffers from similar drawbacks as OTDD: it cannot handle tabular data, images need to have equal resolution and text embeddings need to be of equal size. Moreover, as with OTDD, WTE is only applicable for supervised classification tasks because it explicitly models labels.

The WTE does *not* produce fixed-size dataset characterizations, because the dimensionality of the Wasserstein Embedding is related to the dimensionality of the dataset’s features. Therefore, as mentioned before, it *cannot* have a characterization-based similarity measure by

applying (for instance) cosine similarity on it. However, WTE still supports computing the similarity value with a time complexity constant with respect to the number of datasets by using its inner dataset representation combined with the optimal-transport setup. WTE scales cubically in the number of samples and is thus not scalable with respect to the data. However, since the paper’s experimental evaluation shows the computational cost scaling linearly with the number of datasets, WTE is scalable with respect to the number datasets.

The utility dimension’s aspects are as before due to computing the inner characterization, therefore we only repeat it briefly. WTE cannot be considered a robust similarity measure, because it does not attain invariance properties. Moreover it is debatable whether the produced similarity value is stable. Lastly, the MDS’s dimensionality is considered a parameter.

Comparison and Selection The comparison and selection are guided by the approaches’ aspects, partially depicted in Table 3.1, therewith we can more easily distinguish between approaches. The only aspect not present in the table is the scalability with respect to the number of datasets, which only OTDD cannot attain. Due to the strong connection among the aspects for dataset similarity measures and dataset characterization methods we can re-use the latter’s comparison as part of the current comparison. Therefore, in the context of comparison, we only retain WTE, OTDD and the characterization-based similarity measures for WistubaMF, FeurerMF and Dataset2Vec.

Despite recognizing its great effectiveness potential, we must exclude OTDD because it does not support tabular data well and scales poorly with respect to the number of datasets (and the data). In contrary to dataset characterization methods we do exclude approaches based on scalability, but with respect to the number of datasets. Unlike unscalability with respect to the data, unscalability with respect to the number of datasets limits the suitability of the method in all applicable scenarios –we can adapt the meta-learning approach per dataset, but we do wish to adapt the metadataset, e.g. reducing the number of datasets therein.

Furthermore, we must exclude WTE because it is not applicable to tabular data, does not scale well with the data size and lacks robustness. We do not exclude the remaining approaches since they are scalable with respect to the number of datasets, because they produce fixed-size characterizations allowing for easy (dis)similarity value computation by directly applying cosine similarity or Euclidean distance.

Concluding, the dataset similarity measures to be included in MLTA are the characterization-based similarity measures using the WistubaMF-, FeurerMF- and Dataset2Vec characterization methods. We highlight that the selected collection of approaches is akin to the dataset characterization method selection – all the selected dataset similarity measures are characterization-based measures whose characterization method is incorporated in MLTA – because the aspects we selected them on are strongly connected.

3.1.3 Configuration Characterization Method Selection

In this subsection we select configuration characterization methods for MLTA guided by a comparison on applicability, scalability and utility of the approaches. The materialization of the dimensions (e.g. aspects) for a configuration characterization method is akin to that of a dataset characterization method, but tailored to a configuration instead of a dataset.

Applicability The applicability dimension concerns requirements for the ability to compute a configuration characterization, which depends on its input (the configuration). Recall from

Definition 1 that a pipeline configuration $(g, \mathbf{A}, \boldsymbol{\lambda})$ consists of several components, namely its structure g , algorithms \mathbf{A} and hyperparameters $\boldsymbol{\lambda}$. Regarding the pipeline structure g , we can compare configuration characterization methods on the type of graph they can characterize, with options for a tree with a branching factor of 1 (linear ML pipeline), or a DAG (a pipeline possibly with parallel parts). Regarding the remaining components $\mathbf{A}, \boldsymbol{\lambda}$ we can simply indicate whether the characterization method incorporates them, that is, whether it can distinguish between (some) configurations with varying hyperparameters and/or algorithms. Due to limited knowledge concerning configuration characterizations we do not prefer methods incorporating particular parts thereof; a method not incorporating hyperparameters may still be useful.

Scalability Scalability concerns the ability to perform the characterization of configurations efficiently and in a tractable way. The key aspect here is whether the method scales well with the number of configurations to characterize, because metadatasets may be composed of evaluations for many distinct configurations – after all their search space is infinite. Therefore, we deem a configuration characterization method scalable if its computational time complexity grows linearly with the number of configurations to characterize.

Utility The utility dimension is concerned with the use of a configuration characterization method or its output. Again, we argue for the aspects of parameterization and robustness, but we determine the latter differently. Unlike datasets, configurations do not have meaningless changes associated to them, therefore we are no longer interested in invariance properties. Therefore, we deem a configuration characterization robust if its characterization is stable given the same configuration. Again we need not have deterministic methods, as long as the characterization is stable; presumably a necessary condition to enable a characterization-based meta-learner to effectively use the characterization. At last we again stress the importance of the characterization quality, referring to an effectiveness aspect. However, as before, with a lack of ground truth we can only assess it in a behavioral/experimental way, and do therefore not consider it in the selection of approaches.

RankML [68] Due to its rule-based parsing of the pipeline structure, RankML is able to characterize any pipeline structure g . Moreover, it can distinguish between pipelines with varying algorithms and hyperparameters because it uses a hash-based approach. Furthermore, the setup is sufficiently flexible to also allow for only including algorithms, though this is not the default approach. A drawback of the approach is that it requires the specification of the maximal number of nodes in g for any pipeline in the metadataset. This limitation is however not severe since we typically know which pipelines to characterize a priori to characterizing them. The RankML configuration characterization method is thus applicable on any pipeline $(g, \mathbf{A}, \boldsymbol{\lambda})$.

RankML creates a hash per node in graph g within constant time, therefore the computational time complexity with respect to *one* configuration grows linearly with respect to the number of nodes in g . Because RankML characterizes configurations one by one it also scales linearly with respect to the number of distinct pipelines in the metadataset.

The RankML configuration characterization method is a deterministic approach, given the same collection of pipelines to characterize. In a meta-learning scenario it is safe to assume that the maximal size of the pipelines does not constantly change, therefore we consider RankML to be robust.

Decomposition-Based Configuration Characterizations In this paragraph we consider two approaches simultaneously, namely PMF [45] and TensorOBOE [137], because their approach for pipeline characterization is almost equal, resulting in the same aspects. The beauty and perhaps greatest flaw in using decomposition-based configuration characterizations is its sole dependence on the evaluations, therefore they are applicable on any configuration. Moreover, the methods allow specification of the configuration characterization dimensionality.

Both approaches are not scalable with respect to the number of pipelines to characterize. The simpler matrix-based-decomposition approach is already quite computationally demanding, requiring the decomposition of a high-dimensional matrix with PCA, where the computational cost grows cubically with the number of encountered datasets and quadratically with pipelines (whose configuration space is infinite). With the tensor-based decomposition having worse time complexity it is safe to say that decomposition-based approaches are far from scalable with respect to the number of pipelines to characterize.

Moreover, the sole dependence on the evaluations implies the configuration characterizations depend on the evaluations present for other datasets and configurations. In addition to that, it is questionable whether the characterization is of high quality if pipelines have little evaluations. Furthermore, due to the evaluation dependence characterizations may not be robust to changes in the evaluation collection. Yet, with every evaluation in an AutoML system it changes, and thus each configuration must be re-characterized frequently with a computationally costly decomposition. Another potential cause for instability is that the evaluations of pipelines themselves tend to vary. Concluding, decomposition-based configuration characterizations depend on the other datasets, configurations and the attained evaluation score thereof. Therefore, we do not deem these approaches to be robust, even though we cannot strongly claim the opposite.

Comparison and Selection The comparison and selection of configuration characterization approaches can be brief with only three considered approaches, and two of them sharing aspects due to their great similarity. We exclude the decomposition-based approaches PMF and TensorOBOE from the incorporation in MLTA because they are not scalable with the number of pipelines to characterize. Moreover, we cast doubts on the robustness of these methods. Fortunately, the remaining approach – RankML – attains the desirable aspects and is therefore included in MLTA.

3.1.4 Meta-Learner Selection

In this subsection we select meta-learners for MLTA guided by a comparison on: their applicability regarding knowledge transfer scenarios; their scalability with respect to the metadataset and amount of knowledge transferred; and their utility in terms of parameterization.

Applicability We defer from comparing meta-learners on their applicability with respect to the metadataset, because all discussed approaches are applicable to any metadataset. However, all of them can only incorporate the information from one evaluation metric simultaneously. Nonetheless, we can compare the approaches’ applicability with respect to their knowledge transfer, desiring a broadly applicable approach for both time-restricted (directly retrieving configurations) and warm-starting scenarios.

Scalability Scalability of a meta-learner is mostly concerned with its online phase since this may occur many times. The offline phase may not need full execution on every knowledge transfer because intermediary results can be stored in the AutoML system. Therefore we consider only the online phase scalability, that is when the prior knowledge is transferred to a new dataset. For a meta-learner to be scalable its online phase time complexity should grow at most linearly with the amount of knowledge transferred, e.g. the number of configurations to be recommended or the number of algorithms disregarded from the search space. Moreover, the time-complexity should grow at most linearly with the number of configurations and datasets present in the metadataset.

Utility As with the other meta-learning components we would want the meta-learner to be high-performing, i.e. effective. However, as before we do not compare the approaches on this aspect prior to incorporation because it requires an experimental evaluation. In spite of that, we will assess the parameterization of each approach, preferring approaches that do not require difficult to choose parameters.

Similarity-Based Direct Configuration Transfer This assessment regards the meta-learning approaches [43, 66] that directly transfer configurations from the most similar datasets, either being one configuration or a collection thereof. We can simultaneously assess them because the former approach generalizes the latter. Because these approaches directly transfer configurations they can be used in both time-restricted settings (solely requiring pipelines recommendations) and warm-starting scenarios.

These approaches compute the similarity between the new dataset and any other dataset in the metadataset, considering all of their configuration scores once. Therefore, the time complexity grows linearly with the number of datasets and configurations in the metadataset, and the number of configurations that are transferred – i.e. these approaches are scalable.

The generalization of the approaches requires the specification of the number of datasets to transfer from and the number of configurations to transfer from it. These *two* parameters influence the exploration-exploitation trade-off, where in a warm-starting scenario the optimization budget additionally influences this choice. The prior works have shown that setting both of these parameters to 1 or both to 5 yields fruitful results. Therefore the utility of this approach is debatable, it requires the specification of non-trivial parameters but empirical results suggest promising default values.

Search Space Pruning [94] The approach only allows for pruning search spaces for a subsequent optimization procedure. Therefore it is applicable for warm-starting but not for time-restricted settings, yielding a limited applicability.

This approach assumes a pipeline error represents a sampled error from an algorithm a in pipeline $(g, \mathbf{A}, \boldsymbol{\lambda})$. Then, the offline phase consists of computing the mean-error statistics to create a ranking for the top algorithms, which can be dataset specific. These dataset specific rankings are identified using a similarity value, which needs to be computed for all datasets, for which only the top k components are selected. Therefore, this approach scales linearly with the number of datasets, or constantly in the dataset agnostic setting. The time complexity is constant with respect to the number of configurations and number of components k transferred to the new search space.

The approach requires the specification of parameter k to trade off exploration and exploitation, whose choice may depend on (the size of) the search space, and the subsequent optimization procedure’s computational budget. Ergo, this approach’s utility is limited with a difficult-to-choose parameter.

RankML [68] The meta-learner in the RankML paper is an XGBoost-based ranker model which produces a ranked list of pipelines in the metadataset for the new dataset. Therefore, this approach can both be applied to time-restricted (for instance by selecting the top-ranked pipeline, or the top evaluated pipeline) and warm-starting meta-learning scenarios.

The ranker model is trained in the offline phase to later be used for inference in the online phase. For inference it must compute the dataset characterization for *only* the new dataset. Then, it can predict for each pipeline in the metadataset what its ranking score is. Due to this setup the meta-learner scales well, its online phase time complexity is constant with respect to the number of datasets and evaluations in the metadataset, and linear with respect to the number of distinct pipelines in the metadataset. Concluding, we deem the RankML meta-learner to scale well.

The RankML meta-learner is a parametric approach because it requires the specification of the hyperparameters for the XGBoost-based ranker model. These hyperparameters could be tuned, though this a computationally intensive task. Fortunately, the original paper provides the most important hyperparameters’ tuned values, these can be used as sensible defaults. Unlike the other approaches, the RankML meta-learner does not have parameters that allow for trading off exploration with exploitation. Overall, due to sensible defaults for its parameters, we attribute RankML semi-utility.

Portfolio Building [36] The dataset-agnostic portfolio building meta-learner constructs a portfolio consisting of configurations in its offline phase. The initial interpretation is that this portfolio can be used for warm-starting. However, because the pipelines are added one by one to the collection – based on their reduction of the portfolio’s estimated regret – we can interpret the portfolio as an ordered set, with the configurations ordered by their suitability. This allows for recommending the best configurations directly for time-restricted scenarios. For the aforementioned reasons we conclude that portfolio building attains high applicability.

Due to its dataset agnosticism, portfolio building’s online phase is computationally inexpensive, it simply recommends the previously constructed portfolio or a specified part thereof. Therefore little computation is necessary, yielding constant time complexity with respect to the number of datasets and configurations in the metadataset and the number of configurations transferred. Hence, portfolio building’s online phase is very cheap, computationally speaking, and certainly scalable.

Portfolio building, in its initial interpretation, only needs specification for the portfolio size. This parameter influences the exploration-exploitation trade-off in a warm-starting scenario and may therefore be difficult to select, even though empirical evaluations seem fruitful for a size of 16. However, in a time-restricted setting the parameter is chosen more easily. In a generalized setting we could also parameterize the number of configurations per dataset to consider during portfolio construction. Because, in its initial interpretation, the difficult to select trade-off parameter has a well-performing reference value we deem that portfolio building attains semi-utility.

Comparison and Selection In this paragraph we compare and select the meta-learners on their applicability, scalability and utility. We note that all meta-learners under consideration may have issues with their utility; all but RankML require difficult-to-select parameters to trade-off exploration and exploitation, and RankML itself requires computationally intensive and scenario dependent hyperparameter tuning. Because we cannot exclude all meta-learners we do not exclude any approach based on its utility.

Though all approaches seem to scale well, we note that the dataset-agnostic methods perform best. Accordingly, we do not exclude any meta-learners based on their scalability.

Lastly we consider applicability. We point out that all but the search space pruning meta-learner [94] are widely applicable. Essentially, the search space pruning meta-learner is an antagonist to all other approaches, because it disregards a certain part of the search space instead of promoting it. Therefore, it cannot be used for time-restricted settings requiring configurations to be explicitly formed. Accordingly we exclude the approach from the initial composition of the toolbox. We would like to highlight that future work may still incorporate it, since its application to warm-starting is promising.

Concluding, we incorporate the following meta-learners in MLTA: similarity-based direct configuration transfer [43, 66]; RankML’s meta-learner [68]; and portfolio building [36]. We point out that, according to the previously defined meta-learner categorization (see Section 2.2.2), we yield three differently categorized meta-learners: a similarity-based, a characterization-based and a dataset-agnostic meta-learner. This allows for an interesting comparison of types of meta-learners in an experimental evaluation (see Chapter 4). This is especially important because it is currently unclear which type of approach is to be preferred. For instance, it is not clear whether dataset-agnostic approaches outperform similarity-based approaches due to the findings in the experimental sections of [94, 36] conflicting with each other. The former finds a similarity-based variant outperforms a dataset-agnostic variant of its setup, with the latter concluding the opposite.

3.2 Toolbox Overview

In this section we present the modular framework and functionality of MLTA. In Subsection 3.2.1 we explain the modular framework, detailing the overall structure and its components. In Subsection 3.2.2 we consider the realization of the framework: we discuss the implementation of selected approaches (see Section 3.1) and three additional baselines, perform dataset characterization runtime experiments, and give an overview of MLTA’s provided meta-learning approaches.

3.2.1 Framework

As may have become clear from prior sections, the meta-learning framework consists of the following components: meta-learners, dataset characterization methods, dataset similarity measures, configuration characterization methods, evaluation methods and a metadatabase. The framework is depicted in Figure 3.1, detailing the interactions between components.

From the illustration it is apparent a meta-learning approach is modular, with its required components depending on the type of meta-learner. A similarity-based meta-learner inheriting from `BaseSimilarityLearner` functions with any dataset-similarity measure inheriting from `BaseSimilarityMeasure`, a special case being the `CharacterizationBasedSimilarity` depending on a `BaseDatasetCharacterization` itself. Likewise, any characterization-based

meta-learner inheriting from `BaseCharacterizationLearner` functions with instances inheriting from `BaseConfigurationCharacterization` and `BaseDatasetCharacterization`.

A meta-learner is evaluated using an evaluation method inheriting from `BaseEvaluation`, which interacts with a metadatabase (`MetaDataBase` instance) and a meta-learner (instance inheriting from `BaseLearner`). The interaction with a metadatabase is necessary for accessing the required metadataset, which may be composed of configuration-dataset evaluations but also dataset- and configuration characterizations. Thus, the metadatabase mimics functionality otherwise implemented by an AutoML system, but it also interacts with a `MetaDataTableLookupTable` instance for in-memory look-ups. The tools (implementations inheriting from the aforementioned classes) following this framework are treated in Section 3.2.2, and later used in a warm-starting case study in Chapter 4.

3.2.2 Functionality

3.2.2.1 Component Implementation

In this subsection we discuss details regarding the implementation¹ of previously selected meta-learning components (see Section 3.1) according to the earlier presented framework (see Section 3.2.1). We highlight components whose implementation required the specification of supplementary details. Furthermore, we present two additional meta-learners and one configuration characterization method as baselines incorporated in MLTA.

The Dataset2Vec [58] framework is implemented such that pre-trained networks can be used. The original work, for its cross-validation experimentation, trained five meta-feature extractor networks which we allow to choose as a parameter, with a default for the first trained network. Moreover we allow for specifying the number of batches as a parameter, but with a default value of 10. An example using Dataset2Vec is shown in Figure 3.2.

The RankML [68] configuration characterization method has been implemented following the original work, but we need to explain design decision for unspecified details. We let the hash represent the entire primitive (algorithm including hyperparameters) and not merely the algorithm. The hash is created using the MD5 hashing algorithm [109] on a byte representation – employing UTF-8 encoding [140] – of the full string representation of a scikit-learn [101] pipeline’s step. We use the empty string to represent the ‘blank’ hash and only insert it after other hashes. Given that hash collisions are very unlikely when applying MD5 on the number of distinct primitives in typical metadatasets, these design decisions have as a consequence that every distinct configuration in the metadataset has a unique characterization. Moreover, this interpretation of the framework satisfies its desired deterministic property.

Also the RankML [68] meta-learner needs further specification, especially for incorporation in our modular framework. We use the scikit-learn compatible XGBoost-Ranker implementation from the Python-based `xgboost` package [24] with the approximate trees as specified in [23]. We train this model using the hyperparameter values provided in the paper – using defaults otherwise – on a pairwise ranking objective. For training we assign the ranks $\{0, 1, \dots, N_C\}$ using an evaluation-metric-based ranking, where N_C is the number of configurations used for training.

Furthermore, though not necessary when using the RankML configuration characterization method, we need to specify the behavior when multiple configurations are assigned equal ranking scores, but we cannot recommend all of them. In such a setting we select the configurations

¹See <https://github.com/leightonvg/mlta> for the toolbox.

Meta-Learning Toolbox for AutoML (MLTA) Framework

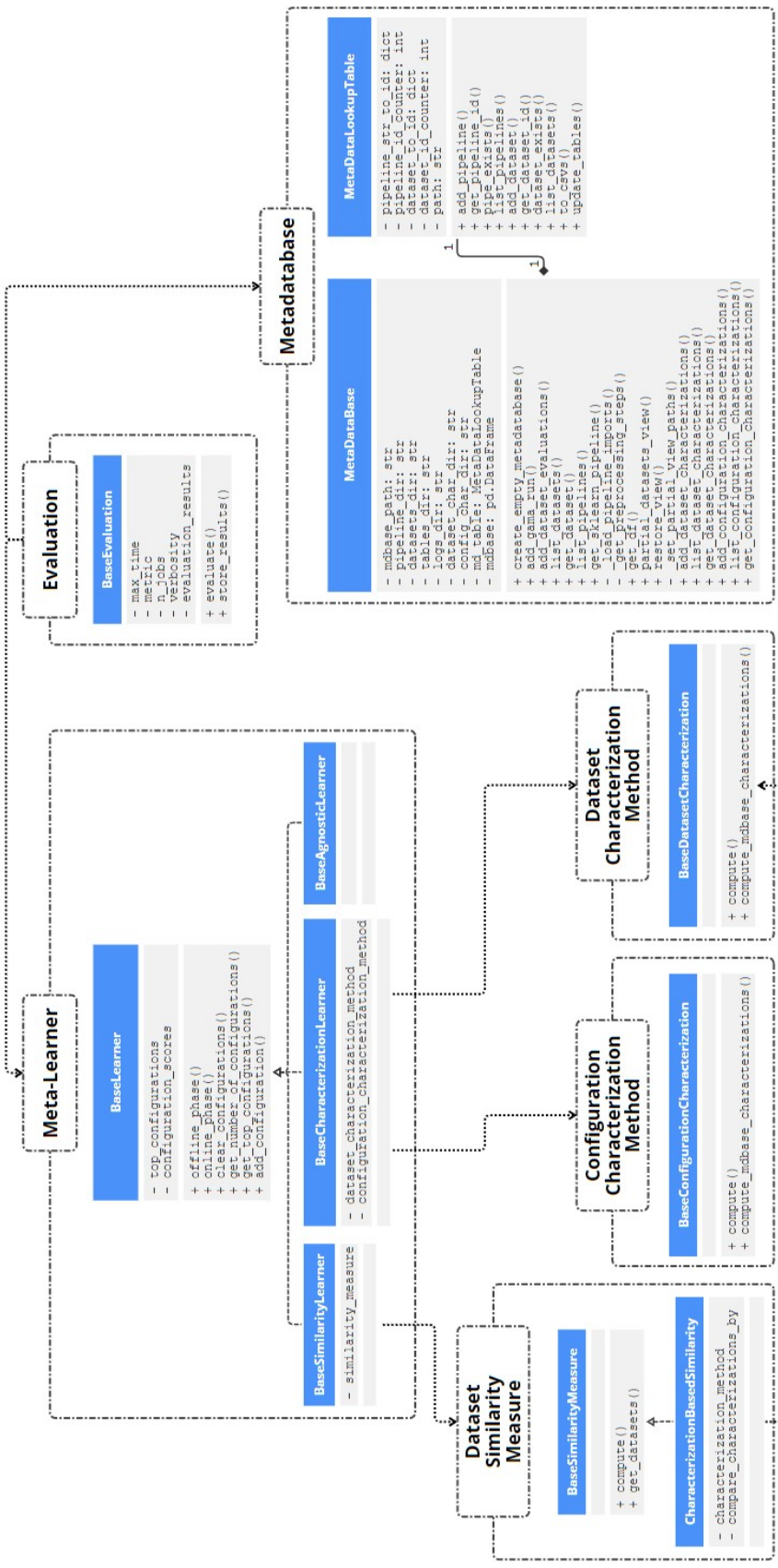


Figure 3.1: A depiction of the MLTA framework, showing the relation between the following main components: meta-learner, dataset-similarity measure, configuration characterization method, dataset characterization method, evaluation and metadatabase.

```

1 from mltा. dataset _characterization import PreTrainedDataset2Vec
2 from mltा. metadatabase import MetaDataBase
3
4 # load metadatabase from "metadatabase_openml18cc" directory
5 mdbase = MetaDataBase(path="metadatabase_openml18cc")
6 # get dataset with id 0 in pd.DataFrame format
7 X, y = mdbase.get_dataset(dataset_id=0, type="dataframe")
8
9 # initialize the characterization method
10 d2v_char_method = PreTrainedDataset2Vec(split_n=0, n_batches=10)
11 # compute the characterization for dataset with id 0
12 d2v_char = d2v_char_method.compute(X, y)
13
14 # compute dataset2vec characterizations for entire mdbase and store them accordingly
15 d2v_chars = d2v_char_method.compute_mdbase_characterizations(mdbase=mdbase)
16 mdbase.add_dataset_characterizations(characterizations=d2v_characters, name="dataset2vec_split0_10batches")
17
18 # retrieve computed characterizations from metadatabase, possibly specifying datasets
19 mdbase.get_dataset_characterizations(characterization_name="dataset2vec_split0_10batches")
20 mdbase.get_dataset_characterizations(characterization_name="dataset2vec_split0_10batches", dataset_ids=[0])

```

Figure 3.2: Code example for dataset characterizations using Dataset2Vec and the metadatabase functionality. The example assumes to have an already created metadatabase in the directory "metadatabase_openml18cc". Other dataset characterization methods operate similarly.

based on their minimal average normalized deviation from the best evaluation score with respect to the dataset the configuration evaluation belongs to. This additional specification is only necessary due to our modular setup; we want to be able to use configuration characterization methods that do not necessarily produce unique characterizations.

We additionally parameterize the specification of the candidate models used during ranker-model inference, and the number of evaluations used during training. The original work used all distinct configurations in the metadataset as candidate models. Accordingly, we set this option as a default, but we allow to specify `max_n_models` – the maximal number of configurations that are considered *per dataset* in the metadatabase, selected on their maximal performance. This option is necessary for reducing the online phase time consumption in time-restricted settings utilizing a vast metadataset. Unlike all other meta-learners RankML did not have parameters for trading off exploration and exploitation, but this is possible with the introduction of `max_n_models`. In Figure 4.1 we give a code example for creating and using the RankML meta-learner to highlight the modular framework of MLTA.

We have implemented a generalization of the dataset-agnostic portfolio building meta-learner [36]. Specifically, we allow for multiple evaluations per dataset in the performance matrix through the `n_pipes` parameter, such that these can be used in the creation of the portfolio. This enlarges the possible portfolio size by, at most, a factor `n_pipes`. Therefore – building on the ordered set interpretation of the portfolio – we allow to specify the number of configurations to be gathered by the online phase's `total_n_configs` parameter, which is upper-bounded by the offline phase's `portfolio_size`.

MLTA provides the implementation of the generalized similarity-based direct configuration transfer approaches as the Top Similarity meta-learner, akin to approaches discussed in Section 2.2.2.1. The Top Similarity meta-learner directly transfers configurations from prior tasks

```

1 from mlta.metadatabase import MetaDataBase
2 from mlta.dataset_characterization import WistubaMetaFeatures
3 from mlta.configuration_characterization import RankMLPipelineRepresentation
4 from mlta.metalearners import TopXGBoostRanked
5
6 # initialize metadatabase from "metadatabase_openml18cc" directory
7 mdbase = MetaDataBase(path="metadatabase_openml18cc")
8
9 # create meta-learner
10 metalearner = TopXGBoostRanked(dataset_characterization_method=WistubaMetaFeatures(),
11                               configuration_characterization_method=RankMLPipelineRepresentation(mdbase=mdbase))
12
13 # offline phase: train the model on entire mdbase but dataset with id 0,
14 # online phase: recommend 25 configurations for dataset with id 0.
15 mdbase.partial_datasets_view(datasets=[0]) # remove all info w.r.t. dataset 0
16 metalearner.offline_phase(mdbase=mdbase)
17 mdbase.restore_view()
18 X, y = mdbase.get_dataset(dataset_id=0, type="dataframe")
19 metalearner.online_phase(X, y, max_time=300, evaluate_recommendations=False, metric="neg_log_loss", total_n_configs=25, max_n_models=500)
20 top25_configs = metalearner.get_top_configurations()
21
22 # 2nd option: use pre-computed dataset & configuration characterizations in mdbase.
23 metalearner.clear_configurations() # remove configurations from prior online phase
24 mdbase.partial_datasets_view(datasets=[0]) # remove all dataset 0 info, also chars
25 metalearner.offline_phase(mdbase=mdbase, dataset_characterizations_name="wistuba_metafeatures",
26                           configuration_characterizations_name="rankml_pipeline_representation")
26 mdbase.restore_view()
27 X, y = mdbase.get_dataset(dataset_id=0, type="dataframe")
28 metalearner.online_phase(X, y, max_time=300, evaluate_recommendations=False, metric="neg_log_loss", total_n_configs=25, max_n_models=500)
29 top25_configs = metalearner.get_top_configurations()

```

Figure 3.3: Code example for a characterization-based meta-learner depicting MLTA’s modularity. It uses the RankML [68] meta-learner, with Wistuba et al. meta-features [134] as dataset characterization method, and the RankML configuration characterization method. For the online phase we specify we desire 25 configurations to be recommended, without evaluating them on the new dataset, in at most 5 minutes time. Additionally, we specified we only want to consider the 500 top-performing configurations per dataset as candidate models. The second example shows how to use pre-computed and stored dataset and configuration characterizations to avoid expensive offline phase computation. The example assumes to have an already created metadatabase in the directory "metadatabase_openml18cc" with dataset and configuration characterizations added. Other meta-learners function similarly.

based on their similarity, where the most similar prior tasks are preferred. To generalize beyond the discussed approaches the Top Similarity parameterizes the number of datasets to transfer from and the number of configurations to transfer from it.

Additional Baselines In addition to the selected meta-learning components (see Section 3.1) we present three baselines: Utility Estimate, a similarity-based meta-learner; Average Regret, a dataset-agnostic meta-learner; and Algorithms Propositionalization, a configuration characterization method.

The similarity-based Utility Estimate meta-learner combines similarity values and configurations’ evaluation scores to determine a configuration’s utility for a new dataset. Firstly,

it computes for each configuration the normalized regret on each dataset it is evaluated on, normalizing with respect to the dataset's evaluation scores. Secondly, it computes the similarity value between the new dataset and all datasets in the metadataset. Then, it computes the configuration utility c_u for each configuration in the metadataset by:

$$c_u = \frac{1}{|\mathcal{D}_{eval}|} \sum_{d=0}^{|\mathcal{D}_{eval}|} ((1 - cr_d) \cdot s_d) \quad (3.1)$$

where \mathcal{D}_{eval} is the set of datasets in the metadataset for which the configuration is evaluated, cr_d is the configuration's normalized regret on dataset $d \in \mathcal{D}_{eval}$, s_d is the similarity value between the new dataset and dataset $d \in \mathcal{D}_{eval}$. Note that $c_u \in [0, 1]$ if $\forall d \in \mathcal{D}_{eval} : s_d \in [0, 1]$, because $cr_d \in [0, 1]$.

After computing the utility scores, the approach recommends the configuration(s) with the highest utility. We deliberately choose a relatively simple approach to act as a baseline for other similarity-based meta-learners. Observe the approach is akin to the similarity-based direct configuration transfer meta-learner (dubbed Top Similarity) and the dataset-agnostic portfolio building meta-learner, because Utility Estimate tries to minimize the regret for the dataset at hand but by taking into account the similarity to that dataset. Unlike portfolio building, it tries to estimate the utility for one configuration on only one dataset, instead of the entire portfolio on all datasets. Unlike Top Similarity, it does not blindly transfer the configurations from a similar dataset, rather it takes their performance into account. A code example for creating the Utility Estimate meta-learner is illustrated in Figure 3.4.

```

1 from mltm.dataset_similarity import CharacterizationSimilarity
2 from mltm.dataset_characterization import FeurerMetaFeatures
3 from mltm.metalearners import UtilityEstimateLearner
4
5 sim_measure = CharacterizationSimilarity(characterization_method=FeurerMetaFeatures(),
6                                         compare_characterizations_by="cosine_similarity")
6 metalearner = UtilityEstimateLearner(similarity_measure=sim_measure)

```

Figure 3.4: Code example for the baseline similarity-based meta-learner Utility Estimate, using a characterization based dataset similarity measure employing Feurer et al.'s meta-features [43] and cosine similarity.

The Average Regret meta-learner acts as a baseline for dataset-agnostic meta-learners. It is similar to Utility Estimate, but it does not require dataset similarity. It computes the normalized regret per configuration in the metadataset, normalized with respect to the dataset's evaluation scores, and ranks the approaches accordingly. In the online phase the highest-ranked configurations with at least `min_evals` evaluations are recommended.

Coming back to the meta-learner comparison's dimensions and accompanying aspects (see Section 3.1.4), both meta-learner baselines are applicable, scalable and have sufficient utility.

At last we present the configuration characterization baseline: Algorithms Propositionalization. This approach encodes the presence of each possible algorithm $a \in \mathcal{A}$ in the configuration, by setting the corresponding entry to 1 if it is present and 0 otherwise. In a similar way it additionally encodes the presence of pairs of algorithms, but in an order sensitive way. These two vectors are concatenated to create the configuration characterization. The size of the configuration characterization is thus $|\mathcal{A}| + |\mathcal{A}|^2$, and hence the characterization tends to be sparse. This method is applicable on any pipeline structure g , scales well with the number of

datasets, configurations and evaluations in the metadataset and is free from parameters. A drawback of the approach is its relatively large dimensionality which depends on the search space. This baseline is deliberately naive in its approach: given a pipeline (g, \mathbf{A}, λ) it only characterizes its algorithms \mathbf{A} , not its accompanying hyperparameters λ nor its *full* structure g . Observe this approach implies the configuration characterizations are not unique.

3.2.2.2 Dataset Characterization Method Run Time Analysis

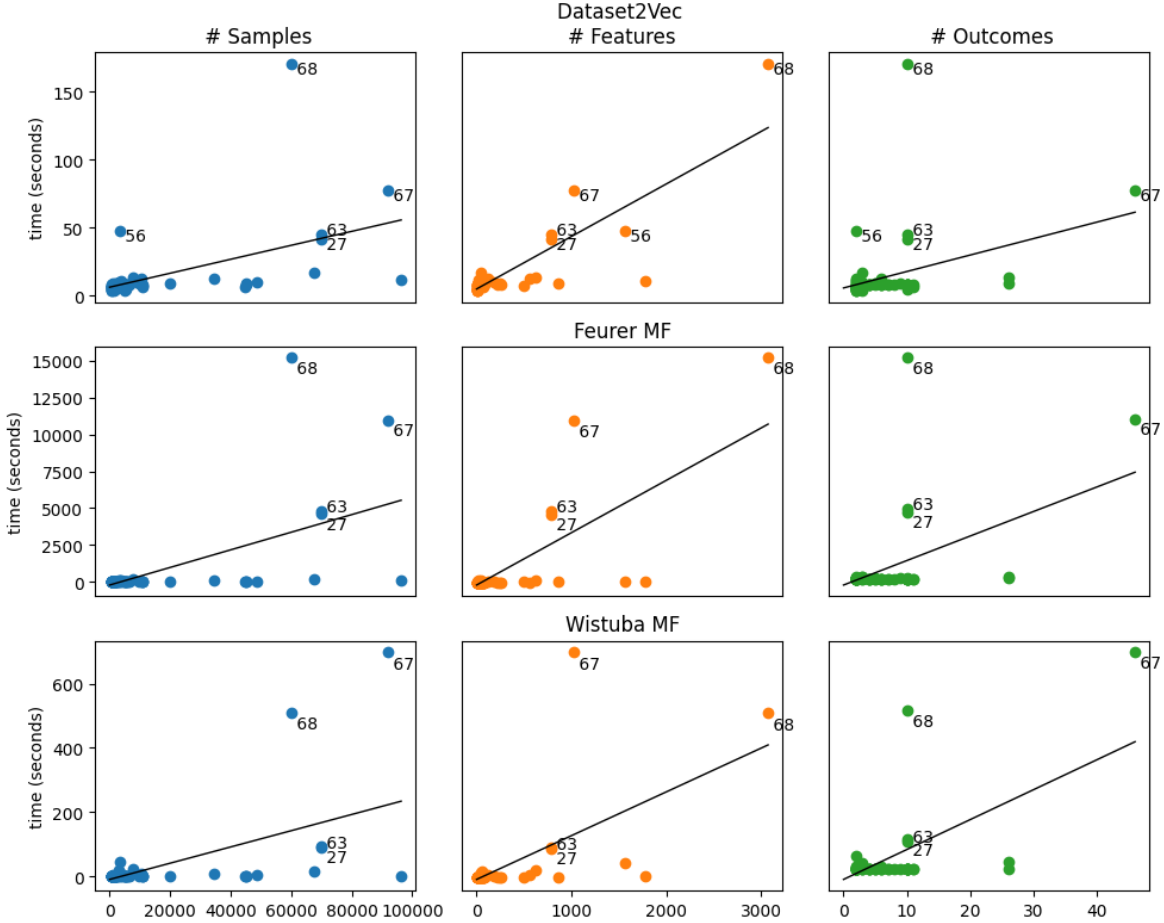


Figure 3.5: Dataset characterization methods' run times plotted against the number of samples, features and outcomes in OpenML18CC [19] datasets. For each cloud of points we depict the best fitting (least-squares) straight line. Outliers are annotated with an id.

In this subsection we analyze the run time of the incorporated dataset characterization methods, hoping to validate the scalability claims in the corresponding selection (see Section 3.1.1). Hence, we analyze the run time behavior for Dataset2Vec [58], Wistuba meta-features [134] and Feurer meta-features [43] with respect to datasets' number of samples, features, and outcomes. We execute the dataset characterization methods using an AMD Ryzen 5 3600XT CPU on OpenML-CC18 [19]: a large ($n=72$), diverse and curated collection of medium-sized classification datasets with extensive use in AutoML-related research. We perform these analyses because the run time of a dataset characterization method has implications for the run

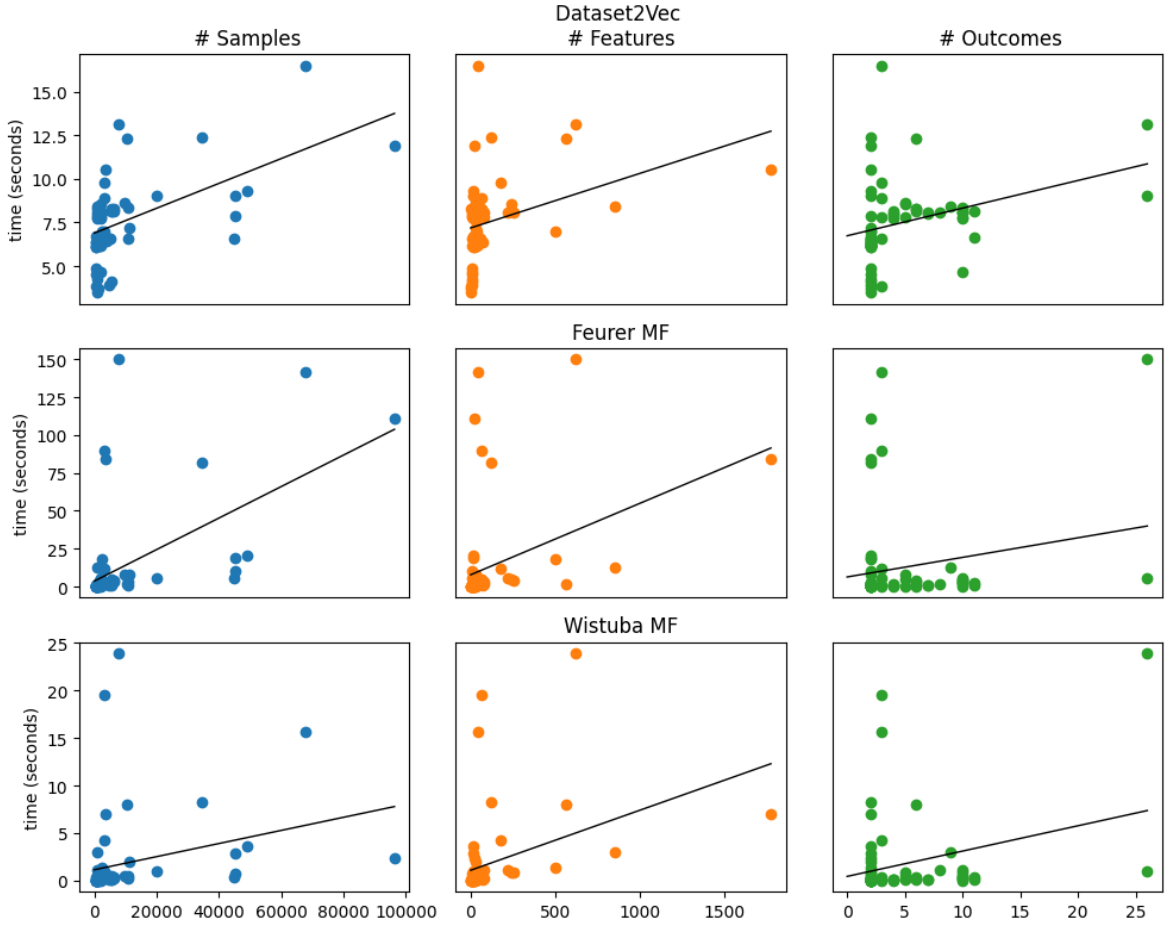


Figure 3.6: Dataset characterization method run times plotted against the number of samples, features and outcomes in OpenML18CC [19] datasets. Similar to Figure 3.5, but excluding its outliers. For each cloud of points we depict the best fitting (least-squares) straight line.

time – and thus viability – of characterization-based meta-learners, but also similarity-based meta-learners employing a characterization-based similarity measure.

The results of this analysis are portrayed in Figure 3.5. None of the scatter plots fit their corresponding line well, rather they violate linear regression assumptions including homoscedasticity and normality of the residuals. Therefore, we cannot confirm the dataset characterization methods scale linearly with respect to the dataset’s properties (number of instances, features and outcomes). The results reveal some datasets require far more time than others, these outliers are annotated and mostly consist of image-based datasets such as MNIST [31] or datasets with lots of features, such as UCI’s Internet-Advertisements [67]. These annotated datasets have many samples, features and/or outcomes, which – especially combined – contribute to increased run time. With exceptions for the outliers, the characterization methods seem to have near constant run time due to the distorted scaling for the y-axis. Therefore, we provide an additional illustration (see Figure 3.6) similar to Figure 3.5 but without the annotated data points, i.e. the outliers. These clouds of points seem to better fit their corresponding least-squares line, but the linear regression assumptions are still violated.

The additional illustration still depicts greater run times for Feurer meta-features, but also reveals only Dataset2Vec requires at least a few seconds for small datasets.

However, we must be careful interpreting these plots because the run time for characterizing a dataset depends on all its properties. Unfortunately, this implies we can not deny nor confirm the aforementioned scalability claims. Additionally, it explains why the height for each dataset’s point is the same across its horizontally neighboring plots. Despite the impossibility of assessing the scalability in absolute fashion, we are able to compare the approaches to each other. Doing so, both figures indicate the Feurer meta-features are more far computationally expensive, with characterization run times up to 4 hours. Furthermore –as exhibited by its greater run time difference between outliers and non-outliers – Feurer MF is less scalable than the other dataset characterization methods, which partially confirms our prior beliefs.

The remaining methods, Dataset2Vec and Wistuba meta-features, seem to be computationally viable, though they also require more time for the annotated datasets. The analysis suggests Dataset2Vec is suitable for datasets with many samples, features and outcomes. Contrarily, the Feurer meta-features are not suitable for larger datasets. The Wistuba meta-features are far less computationally expensive than the Feurer meta-features, it only struggles with (the two largest) datasets simultaneously having many samples, features, and outcomes.

Additionally, we want to point out that the datasets annotated with id 63 and 27 possess similar properties and exhibit similar behavior, in fact these are the intimately connected MNIST [31] and Fashion-MNIST [136] datasets. Though not the initial intent of the analysis, we do highlight this finding suggests all considered dataset characterization methods are stable in terms of their run time.

3.2.2.3 Provided Meta-Learning Approaches

In this subsection we briefly highlight the power of MLTA’s modularity. In Figure 3.2 we give an overview of all meta-learning approaches that can be constructed using MLTA’s provided meta-learning components. With 5 meta-learners we can create 14 meta-learning approaches using 2 configuration characterization methods, and 3 dataset characterization methods with their accompanying characterization-based similarity measures. Moreover, we can alter the approaches’ parameters, such as using Euclidean distance instead of cosine similarity for characterization-based similarity measures. In addition to that, all meta-learners are implemented to have the possibility to evaluate configurations in the online phase. By doing so they return only a selection of the top-performing configurations. We have previously demonstrated that creating these meta-learning approaches requires only few lines of code, see Figures 3.2, 4.1 and 3.4.

Keep in mind – as shown in the dataset characterization method run time experiments – approaches’ suitability may vary depending on the task. Unfortunately, we cannot replicate the implemented methods’ performances because they are dependent on the original works’ metadatasets, which are not made publicly available. However, we are eager to compare these meta-learning approaches on their effectiveness in an experimental setting (see Section 4.2), and evaluate their use for incorporation in an AutoML system (see Section 4.3).

Meta-Learning Approach	Meta-Learner	Dataset-Similarity Measure	Dataset Characterization	Configuration Characterization
AverageRegret	AverageRegret			
PortfolioBuilding	PortfolioBuilding			
UtilityEstimate-Wistuba	UtilityEstimate	Wistuba Cosine Similarity		
UtilityEstimate-Feurer	UtilityEstimate	Feurer Cosine Similarity		
UtilityEstimate-Dataset2Vec	UtilityEstimate	Dataset2Vec Cosine Similarity		
TopSimilarity-Wistuba	TopSimilarity	Wistuba Cosine Similarity		
TopSimilarity-Feurer	TopSimilarity	Feurer Cosine Similarity		
TopSimilarity-Dataset2Vec	TopSimilarity	Dataset2Vec Cosine Similarity		
RankML-Wistuba	XGBoost-Ranker		Wistuba Meta-Features	RankML
RankML-Feurer	XGBoost-Ranker		Feurer Meta-Features	RankML
RankML-Dataset2Vec	XGBoost-Ranker		Dataset2Vec	RankML
AP-Wistuba	XGBoost-Ranker		Wistuba Meta-Features	Algorithms Propositionalization
AP-Feurer	XGBoost-Ranker		Feurer Meta-Features	Algorithms Propositionalization
AP-Dataset2Vec	XGBoost-Ranker		Dataset2Vec	Algorithms Propositionalization

Table 3.2: Overview of all meta-learning approaches we can create using tools provided by the modularized MLTA framework. The columns Meta-Learner, Dataset-Similarity Measure, Dataset Characterization and Configuration Characterization indicate the presence of the incorporated meta-learning components (see Section 3.1 and 3.2.2.1), with blanks indicating absence thereof. The meta-learning approaches are grouped by their meta-learner categorization, the first two approaches are dataset-agnostic, the following six are similarity-based, and the last six are characterization-based meta-learning approaches.

Chapter 4

MLTA Case Study: Meta-Learning for Warm-Starting Pipeline Search

In the previous chapter (3) we introduced the Meta-Learning Toolbox for AutoML (MLTA), in the current chapter we display its utility. In particular, we explore the use of MLTA for warm-starting evolutionary pipeline search, as provided by the AutoML research framework GAMA [47]. Thereby we aim to validate our research objective: create a toolbox to easily construct, analyze and compare existing and novel meta-learning approaches and components.

As mentioned previously (see Section 2.2.2.1), the goal of warm-starting is to attain similar performance at reduced computational cost, or increased performance at similar computational cost, or preferably increased performance at reduced computational cost. To allow for assessing these goals and our research objective we execute the case study in a 2-step approach. We additionally aim to mimic the real-life usage as a prototyping tool, where limited computational resources prohibit the enumeration of all options for computationally demanding experimental evaluations – e.g. warm-starting GAMA.

In the first step we construct MLTA’s meta-learning approaches to compare and analyze them in a time-restricted setting. Therein we grant only 5 minutes of online learning to recommend 25 previously-seen pipeline configurations, thus *without* tuning configurations – we simply transfer and evaluate them. These results are used to identify the meta-learning approach with the highest potential for warm-starting, without actually performing warm-starting. This allows us to perform the second step: incorporate and evaluate the highest-potential meta-learning approach for warm-starting GAMA.

We stress the goal of this case study is *not* necessarily to improve GAMA’s search with warm-starting. Instead, by performing the 2-step case-study we aim to validate our research objective, justifying the claim that we created a toolbox that allows for easily constructing, analyzing and comparing existing and novel meta-learning approaches and components.

The remainder of this chapter is structured as follows. In Section 4.1 we discuss the creation of a set of previous experiences – a metadataset – to be used by the meta-learners, using the metadatabase functionality provided by MLTA. In Section 4.2 we discuss the first step of the case study: an analysis of meta-learning approaches with a time-restricted online phase, therein we employ meta-learning approaches facilitated by MLTA. Using the results from the first step, we explore and evaluate warm-starting evolutionary pipeline search in Section 4.3.

4.1 Metadataset

A large part of creating a meta model is, as with any ML model, the data itself. Many existing works create a metadataset specifically for their own purpose, which is rarely shared. Hence, we need to create a high-quality metadataset ourselves for comparing meta-learning approaches, ideally in a broad manner allowing others to reuse it.

A fully complete metadataset would contain lots of information by exhaustively evaluating each solution in the configuration search space, possibly multiple times for solutions with varying performance, all on many different datasets and performance metrics. It quickly becomes apparent that creating a fully complete metadataset is computationally infeasible. Even more so, in our AutoML formalization (see Definition 1) we do not restrict the size of pipelines, effectively allowing for an infinite pipeline length and hence infinite search space. Fortunately, recent work suggests an opportunistic, and possibly unreliable, set of prior experiences can still be fruitful for guiding AutoML search [94]. Though we cannot and need not construct an exhaustive metadataset, we still care about completeness because it is directly related to the external validity of our case study.

A main consideration for any metadataset is the collection of datasets – or tasks – to consider, aligning with MLTA’s focus we only consider classification datasets. For the creation of the metadataset we consider OpenML-CC18 [19]: a large ($n=72$), diverse and curated collection of medium-sized classification datasets which has been extensively used in AutoML-related research. The next main consideration is which configurations’ performance we want to record and on which of the datasets in OpenML-CC18.

This choice is particularly difficult due to an infinite number of configurations and because we need to trade off the metadataset’s completeness with its configurations’ performance, which could be dataset dependent. We propose to overcome this difficulty by deferring the choice to AutoML pipeline search, recording the dataset-configuration evaluations it encounters in the process. Pertaining to metadataset creation, existing search methods differ on two main aspects: completeness and performance, which we can trade off using hyperparameters. But, computational cost prohibits the enumeration all possible search methods and their parameters. Rather, to trade off the metadataset completeness and its configurations’ performance we propose to use multiple distinct search methods – varying on their completeness-performance trade-off – to populate the metadataset with prior experiences, namely:

- Random Search: Randomly pick machine learning pipelines from the search space and evaluate them.
- Asynchronous Evolutionary Algorithm: Evolve a population of machine learning pipelines, drawing new machine learning pipelines from the best of the population.
- Asynchronous Successive Halving Algorithm: A bandit-based approach where many machine learning pipelines iteratively get evaluated and eliminated on bigger fractions of the data.

with random search focusing on completeness, the Asynchronous Successive Halving Algorithm being performance-oriented, and the Asynchronous Evolutionary Algorithm focusing on both aspects. Note, the latter two algorithms depend on their initialization and could benefit from multiple runs but in the creation of the metadataset we only employ them once, arguing the composition of multiple search methods yields sufficient quality, especially considering multiple runs may yield similar results and thus waste limited computational resources.

The metadataset quality depends on the performance of the configurations found during the search methods, whose performance is mainly impacted by the amount of resources dedicated to optimization. The aforementioned search methods to populate the metadataset are incorporated in GAMA [47], which has been shown to achieve satiable performance on the OpenML-18CC datasets given an optimization budget of 1 hour, 32GB of RAM and 8 vCPUs [46]. To create the metadataset we employ GAMA with the aforementioned search methods and optimization budget for each search method separately, using otherwise default hyperparameters. We store these results using MLTA’s `MetaDataSet` functionality.

However, we store encountered configurations and their associated negative logistic loss performance only if they are evaluated on a full dataset, thus neglecting many evaluations during the Asynchronous Successive Halving Algorithm. The stored performance estimate for a pipeline configuration is reliable, because it is computed using 5-fold cross-validation. Due to employing multiple search methods, the metadataset may contain multiple evaluations of the same configuration on the same dataset using the same metric, we aggregate these instances by averaging their negative logistic loss score.

Performing the aforementioned process required 216 hours of computational time – with the aforementioned optimization budget – to yield a metadataset composed of 985.475 configuration evaluations and 826.270 distinct pipeline configurations. Though each configuration is evaluated ≈ 1.2 times on average, there are plenty of configurations with numerous evaluations since the distribution is extremely long tailed. Only 66.050 pipelines are evaluated more than once totalling 225.255 evaluations. In similar fashion, the metadataset is composed of 534 configurations with at least 25 evaluations. The number of evaluations per dataset varies greatly with the most and least evaluated dataset having 76.945 and 21 evaluations respectively (see Appendix B), which varies due to differences in dataset size and associated configuration evaluation time. All non-image datasets in OpenML-18CC have at least 400, but often far more, configuration evaluations. We include these image-based datasets because they can be interpreted in a tabular format, and because they are part of OpenML-18CC, which we do not want to alter unnecessarily.

Another approach for creating a metadataset, requiring no to little compute resources, is to use OpenML[127]. OpenML contains many evaluations (runs) and pipeline configurations (flows), but due to some issues it is not well suited as a source for creating our metadataset:

- The flows, and associated runs, are spread across distinct machine learning frameworks, with varying versions themselves. Because performance may vary across implementations and versions, combining these results is non-trivial.
- With complete ML pipelines in mind, many datasets have an underexplored search space, which may lead to a poor-quality metadataset in terms of completeness. Typically, preprocessing transformations are simple, if there are any. Moreover, apart from a bias towards pipeline structure, there often is a bias towards certain aspects of the pipeline. For instance, there are users contributing many evaluations only exploring the effect of hyperparameters, disregarding algorithm selection for preprocessing and prediction.

Co-occurrence of the aforementioned issues further restrict OpenML’s suitability as a source for our metadataset. Therefore, despite containing numerous previous experiences, OpenML is poorly suited for directly creating the metadataset we desire.

4.2 Time-Restricted Meta-Learning

In this section we use the introduced metadataset (Section 4.1) as previous knowledge for meta-learners whose online phase is time restricted. Evaluating meta-learners in this time-restricted setting has three goals; firstly it aims to compare and analyze meta-learners’ performance when employed with limited computational resources; secondly it aims to identify the highest potential meta-learner to later be used in warm-starting pipeline search (see Section 4.3); thirdly it serves as a validation for the research objective: a toolbox to easily construct, analyze and compare existing and novel meta-learning approaches and components.

We construct all meta-learning approaches provided by MLTA’s modularized framework (see Table 3.2). For each meta-learning approach we perform a leave-one-dataset-out evaluation on OpenML-CC18: we evaluate each meta-learner on each dataset in OpenML-18CC, using the remaining datasets’ metadata as previous experience to learn from. Each meta-learner is given 300 seconds of online phase computation time, and unlimited offline phase time. In the online phase the meta-learner is tasked with recommending 25 pipeline configurations suitable for the left out dataset. The meta-learner does not get access to the dataset and associated metadata it is evaluated on. Nor does it get the possibility to evaluate and tune the recommended evaluations on the new dataset, because that is the job of a subsequent optimization procedure which we do not yet consider. We evaluate the 25 recommended pipeline configurations on negative logistic loss using a 80/20 holdout split on the left out dataset.

We evaluate meta-learners using the MLTA framework (see Figure 3.1). We combine instances indirectly inheriting from `BaseLearner` – the meta-learners – with an evaluation method inheriting from `BaseEvaluation` – a `LeaveOneDatasetOutEvaluation` – and a `MetaDataBase` instance giving access to the metadataset and providing AutoML functionality. Figure 4.1 shows that we can construct, execute and evaluate a meta-learner with few lines of code.

```

1 from mlta.metadatabase import MetaDataBase
2 from mlta.metalearners import PortfolioBuilding
3 from mlta.evaluation import LeaveOneDatasetOutEvaluation
4
5 mdbase = MetaDataBase(path="metadatabase_openml18cc")
6 metalearner = PortfolioBuilding()
7 evaluation_method = LeaveOneDatasetOutEvaluation(validation_strategy="holdout",
8     test_size=0.2, n_configs=25, max_time=300)
evaluation_results = evaluation_method.evaluate(mdbase, metalearner)
```

Figure 4.1: Code example for evaluating a meta-learner. Due to the MLTA framework we need not bother with evaluation, dataset and metadataset related operations. The example assumes to have an already created metadatabase in the directory "metadatabase_openml18cc". Other meta-learners can be evaluated similarly.

We compare the performance of meta-learning approaches by the negative logistic loss of the their highest-ranked pipeline configuration recommendation on each dataset and rank the approaches accordingly. To mimic real-life use in a time-restricted setting we select the highest-ranked configuration by only allowing a handful of evaluations. Specifically, we evaluate the 5 configurations ranked highest by the meta-learner and use the negative logistic loss from the top-evaluated pipeline (among the 5 pipelines) to create the ranking. Not only do we believe this evaluation procedure mimics real-life usage, we believe it may achieve satiable performance; research suggests selecting the meta-learner’s best-evaluated pipeline from a handful of its top

recommendations yields comparable performance to AutoML [68].

To ground the meta-learner results we include the three one-hour pipeline optimization results from the metadataset (see Section 4.1) utilizing Random Search, the Asynchronous Evolutionary Algorithm (AsyncEA) and Asynchronous Successive Halving Algorithm (ASHA), using their best-evaluated pipeline configuration. Note that the scores for these reference AutoML results are computed using a 5-fold cross-validation and are therefore comparable to the 80/20 hold-out split, though the reliability of the estimates differ. Since the meta-learners do not get the possibility to evaluate and tune the recommended configurations on the new dataset, it would be surprising – but possible – if they outperform the 1-hour AutoML reference scores with only 5 minutes of online phase time.

Recall from Chapter 3 that MLTA allows the pipeline configuration to be evaluated prior to recommending them, since it is a possibility for any of the discussed meta-learners. Because we are interested in warm-starting pipeline optimization we do not evaluate configurations before recommending them, because they will be evaluated within the optimization procedure itself. It is unknown whether evaluating configurations before recommending them for warm-starting pipeline optimization is beneficial, investigating this is out-of-scope for this thesis, but it could be a topic for follow-up investigation.

As previously mentioned, we aim to mimic real-life usage of the toolbox where computational resources are not abundantly available. Yet, this aim does not prohibit the enumeration of all meta-learning approaches in this initial analysis, because each approach is time constrained.

Note that despite the time-restricted setting we grant the meta-learner unlimited time during the offline phase because this prior information can be pre-computed and accumulates over time, where AutoML systems can often avoid re-computation, in turn yielding a relatively inexpensive offline phase. For instance, to employ a characterization-based similarity measure we need not compute characterizations for prior tasks, only for the new task, which should be inexpensive compared to the online phase and a potential subsequent pipeline optimization procedure.

The current setup requires $72 \cdot 14 = 1008$ dataset-meta-learner combinations to be evaluated, each requiring their own offline phase, online phase and evaluation phase. In other words we need to do the following 1008 times: construct a meta-learner, use it for recommending 25 pipelines and evaluate each of them. Therein the typical parallelization while using cross-validation is infeasible, because the meta-learners themselves are already allotted the previously mentioned computational resources (8 vCPUs and 32GB of RAM). Therefore we employ holdout during the leave-one-dataset-out evaluation procedure.

Unfortunately, with the aforesaid setup we are unable to adequately assess all approaches due to a data leakage issue: Dataset2Vec employs a pre-trained network trained on 120 UCI classification datasets, with some of them occurring in OpenML-CC18. Despite that Dataset2Vec was trained on a different task (dataset similarity prediction) we cannot evaluate it on OpenML-CC18, because its pre-trained network has seen parts of the supposed-to-be test set. We have two options: either exclude UtilityEstimate-Dataset2Vec, TopSimilarity-Dataset2Vec, RankML-Dataset2Vec and AP-Dataset2Vec from the analysis, or alter OpenML-CC18. Instead of preferring one analysis over the other, we provide the results for both analyses in Section 4.2.1. Thus, we provide the experimental results for the full OpenML-CC18 collection, which excludes the Dataset2Vec approaches. In addition to that, we provide the results for an OpenML-CC18 subset – a selection of 38 out of 72 datasets – excluding Dataset2Vec’s pre-training datasets (see Appendix D). Beware that this selection does *not* imply a different metadataset.

4.2.1 Results

Ranks We aim to rigorously compare the ranks using a critical differences (CD) diagram as introduced by [29]; a CD diagram is a compact visual representation of the approaches' average ranks and significance tests on pairs of approaches with horizontal lines grouping statistically indistinguishable approaches. However, before we are able to produce a CD diagram we must first test whether there is any difference among the meta-learning approaches, for which we employ the non-parametric Friedman test [44]. For both the full OpenML-CC18 collection and the subset we can reject the Friedman test's null hypothesis with $p < 0.0001$ and $p \approx 0.0107$ respectively. We depict the CD diagrams comparing the meta-learning approaches and the pipeline optimization procedures in Figures 4.2 and 4.3. Therein we do not employ Nemenyi's test as proposed in [29], rather we use a Wilcoxon signed-rank test since it is favored by more recent work [16]. In employing these tests we consider a significance level $\alpha = 5\%$, applying a Bonferroni correction [116] to correct for multiple testing.

The CD diagram for the OpenML-CC18 subset (Figure 4.2) reveals that we cannot reject equivalence of any meta-learning approach with any baseline. Please note that this *does not imply these approaches perform equivalently* because "absence of evidence is not evidence of absence" [10]. Perhaps surprisingly, we can claim that Portfolio Building outperforms 5 other meta-learning approaches, namely: AP-D2V, AP-Wistuba, UtilityEstimate-Feurer, TopSimilarity-Feurer and RankML-Wistuba. Thus, in context of meta-learner categorization, the dataset-agnostic portfolio building outperforms three characterization-based meta-learners and two similarity-based meta-learners.

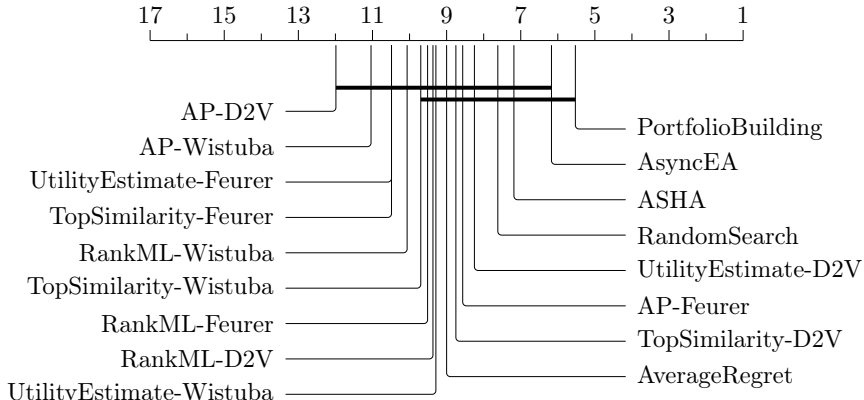


Figure 4.2: Critical differences (CD) diagram comparing meta-learning approaches on the OpenML-CC18 subset. We can create this CD diagram a – i.e. perform the pairwise tests – because we reject the null hypothesis for the corresponding Friedman test ($p \approx 0.0107$).

The CD diagram for the full OpenML-CC18 collection (Figure 4.3) tells a similar story. Compared to the OpenML-CC18 subset we now have more measurements – which increases the power of our statistical tests – resulting in more statistically significant distinctions among approaches. Now we can confirm that the baselines outperform a meta-learner, namely AP-Wistuba. It indicates that the combination of a characterization-based meta-learner with simple dataset- and configuration characterizations may not be the best approach for knowledge transfer. Moreover, this analysis confirms that Portfolio Building is the best meta-learner: with statistical significance we can state that Portfolio Building outperforms all other meta-learners.

Yet, we cannot reject Portfolio Building's equivalence from the metadataset baselines. Again we emphasize that "absence of evidence is not evidence of absence" [10], and these results do therefore not mean that Portfolio Building performs equivalently to the metadataset baselines.

Despite varying average ranks, we have found little statistically significant differences between meta-learning approaches. Therefore we can not claim that certain meta-learning *components* outperform others.

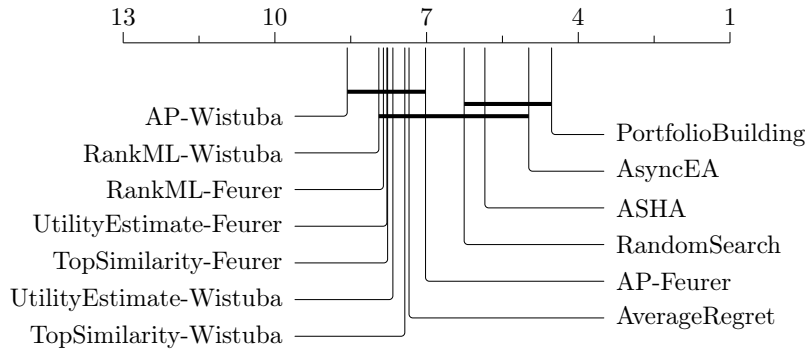


Figure 4.3: Critical differences (CD) diagram comparing meta-learning approaches on the full OpenML-CC18 collection. It excludes the Dataset2Vec-based approaches due to aforementioned data leakage issues. We can create this CD diagram – i.e. perform the pairwise tests – because we reject the null hypothesis for the corresponding Friedman test ($p < 0.0001$).

We visualize the rank distribution for each approach in Figures 4.4 and 4.5. Beware that these illustrations do not allow for well-grounded comparisons on average ranks. Rather, they allow for additional qualitative analyses of the rank distributions.

We observe that the approaches' rank distributions are comparable for both figures, i.e. their kernel densities are similar in shape. The shape thereof indicates the approaches' consistency, with more uniform densities indicating less consistency. On this note, we observe some approaches to more consistently rank poorly, including AP-D2V, AP-Wistuba, UtilityEstimate-Feurer, TopSimilarity-Feurer, TopSimilarity-Wistuba – a list very comparable to the approaches outperformed by PortfolioBuilding for the OpenML-CC18 subset (Figure 4.2). Contrarily, both AsyncEA and PortfolioBuilding more consistently receive high ranks. Notably, this consistency is greater for PortfolioBuilding than for any other meta-learning approach, and perhaps some of the baselines. Lastly, both Figures indicate that AsyncEA seems to be more consistent in achieving better ranks than the other baselines.

Considering the OpenML-CC18 subset (Figure 4.4), we note that none of the metadataset baselines receive the two worst ranks among any dataset. Moreover, AsyncEA never obtains the 3 worst ranks and RandomSearch never attains the best ranking. This behavior is only partially visible for the full OpenML-CC18 collection (Figure 4.5), there only AsyncEA never receives the worst ranking.

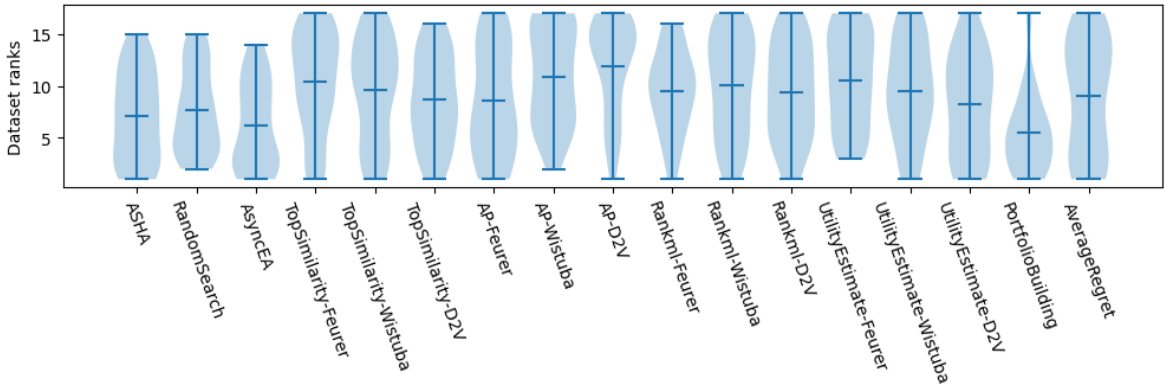


Figure 4.4: Violin plot of the dataset ranks attained on the OpenML-CC18 subset.

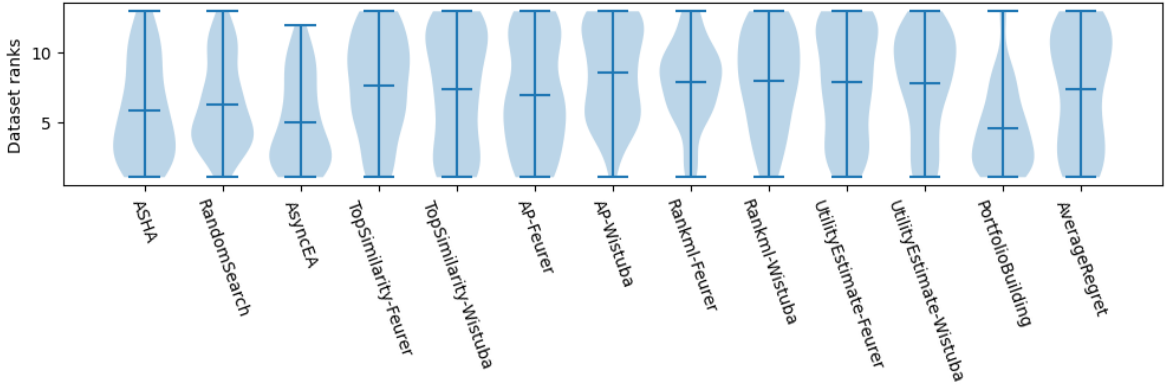


Figure 4.5: Violin plot of the dataset ranks attained on the full OpenML-CC18 collection.

Regret Selecting the meta-learning approach with the highest warm-starting potential is not as simple as taking the method with the highest average rank in the CD diagram, because it only considers the top-ranked recommended pipeline configurations. For warm-starting we provide a larger set of configurations, thus a meta-learner selection could account for it. We are unaware of any work exploring warm-starting potential, that is, extrapolating a meta-learning approach’s evaluation results in a time-restricted setting to the final performance obtained by using it for warm-starting AutoML search.

We are not attempting to create such a score for it heavily relies on the exploration-exploration trade off. Rather, we aim to gain better insights into the performance of the proposed approaches by providing their aggregated regret and standard deviation in Table 4.1. In essence the aggregated regret is the average normalized regret with respect to both datasets and its recommended pipeline configurations, a formalization can be found in Appendix E. The aggregated standard deviation is the standard deviation of the recommendations’ evaluation scores, averaged over datasets.

In the context of warm-starting, we consider the aggregated regret to be a proxy for the exploitation focus of a meta-learning approach; a lower regret prioritizes higher evaluation scores and punishes for missing recommendations. Contrarily, the standard deviation is a proxy for the diversity – or exploration in a warm-starting context – of the meta-learning

approach. These values provide an additional – but not leading – analysis into the exploration and exploitation of the presented meta-learning approaches. Note that the regret values are informative in a relative sense instead of absolute; they depend on the performance of the other approaches.

Meta-Learning Approach	Regret Full OpenML-18CC	SD Full OpenML-18CC	Regret Subset OpenML-18CC	SD Subset OpenML-18CC
AverageRegret	0.1012	0.0992	0.1366	0.1295
Portfolio Building	0.1462	0.2041	0.1919	0.2437
UtilityEstimate-Wistuba	0.1484	0.0941	0.2116	0.1165
UtilityEstimate-Feurer	0.1722	0.0997	0.2338	0.1008
UtilityEstimate-Dataset2Vec			0.1975	0.1452
TopSimilarity-Wistuba	0.1559	0.1547	0.2119	0.1851
TopSimilarity-Feurer	0.1601	0.1376	0.2325	0.1601
TopSimilarity-Dataset2Vec			0.1822	0.1662
RankML-Wistuba	0.1761	0.1861	0.2286	0.2093
RankML-Feurer	0.2108	0.1942	0.2817	0.2002
RankML-Dataset2Vec			0.2113	0.2193
AP-Wistuba	0.2146	0.0615	0.2970	0.0770
AP-Feurer	0.1684	0.0564	0.2268	0.0398
AP-Dataset2Vec			0.1687	0.0860

Table 4.1: Aggregated regret and standard deviation for meta-learning approaches on the OpenML-CC18 collection [19] and its subset (Appendix D). The regret is computed according to Equation E.2.

We note that the AverageRegret meta-learner achieves the lowest regret for both the full OpenML-18CC collection and its subset, we claim this is because it recommends configurations based on the aggregated regret. It is a prime example that a lower aggregated regret does not imply a high average rank in the CD diagrams. For warm-starting evolutionary search we seek a meta-learner that has relatively low regret but a relatively high amount of standard deviation.

We remark that all meta-learning approaches including a baseline meta-learning component (Algorithms Propositionalization, UtilityEstimate and AverageRegret) achieve low standard deviation in their recommendations. We hypothesize that the approaches including algorithms propositionalization achieve low variety because the configuration characterization method cannot distinguish well between configurations with varying hyperparameters and pipeline structure. Due to this incapability it is likely many similar pipelines are recommended, which in turn obtain similar evaluation scores. Thus, it seems that for warm-starting AutoML we prefer configuration characterization methods that can distinguish pipelines with varying hyperparameters and structures well. We are not certain why the AverageRegret and UtilityEstimate and approaches obtain low standard deviation. A possibility for the UtilityEstimate meta-learners is that the evaluation scores for the most similar dataset are very close to each other. Then, the meta-learner could completely disregard other datasets and only sample configurations from one dataset, whose top configurations are likely similar.

Out of all approaches Portfolio building has the highest standard deviation for both the full OpenML-18CC collection and its subset, while maintaining relatively low levels of regret. In fact, there is no approach that simultaneously has lower regret and higher standard deviation for both OpenML-18CC variants. We suppose Portfolio Building’s high configuration diversity is achieved by only considering the top performing configuration per dataset, thereby sampling configurations from 25 distinct datasets. The other meta-learning approaches also consider the configurations that did not achieve the best performance on the dataset, and are thus more likely to include configurations from a smaller number of datasets.

Selecting a warm-starting approach We should not select a meta-learning approach for warm-starting purely on one aspect. In that spirit, we compared approaches on their average rank, rank distributions, aggregated regret and standard deviation. Portfolio building outperforms (all) other meta-learning approaches on their average dataset ranks, achieves consistent low ranks on each of the datasets, has relatively low aggregated regret and high configuration diversity – the latter being especially important for warm-starting evolutionary search. Therefore, we conclude that Portfolio Building is the meta-learning approach with highest potential for warm-starting AutoML search. Though we cannot make strong claims, the analyses seem to indicate that AsyncEA is the highest-performing search method. Therefore we warm-start AsyncEA using Portfolio Building in Section 4.3.

4.3 Warm-Starting Asynchronous Evolutionary Pipeline Search

In this section we aim to further validate our research objective – the creation of a toolbox to easily construct, analyze and compare existing and novel meta-learning approaches and components – by using MLTA for warm-starting AutoML search. However – unlike the previous section – we do not compare meta-learning approaches and components, because warm-starting experimentation requires far more computational resources. In the previous section we have selected the portfolio building meta-learning approach for warm-starting asynchronous evolutionary pipeline search using GAMA, which we will refer to as warm-started AsyncEA from now on. We note that we could warm-start the asynchronous successive halving algorithm in similar fashion, but we cannot warm-start random search.

We allot warm-started AsyncEA one hour of total computation time for pipeline optimization, that is excluding time spent in metadataset related operations such as data loading. Therein we allow the meta-learner to use a maximum of 5 minutes for recommending at most 25 pipeline configurations to warm-start AsyncEA, the remaining computation budget (at least 55 minutes) is spent on the pipeline search. The (at most 25) recommended pipeline configurations are incorporated in the initial population consisting of 50 individuals in total, with the remaining (at least 25) individuals being generated randomly.

We have no particular reason to choose for 5 minutes of meta-learning computation time. To the best of our knowledge there are no works that could indicate sensible values, though amounts larger than the optimization budget are clairvoyantly nonsensical. For most meta-learning approaches the amount of time may be a crucial aspect, because it influences the exploration exploitation trade-off for the subsequent warm-started pipeline optimization. For the current dataset-agnostic portfolio building approach we expect the online phase to almost immediately return the warm-starting individuals – in substantially less than 5 minutes – because we only need to recommend the configurations identified in the offline phase. Therefore,

we do not expect the results *for this meta-learning approach* to depend much on the amount of time allotted to it.

We opt for an initial population totalling 50 individuals because that is the default value provided by GAMA. We suppose that replacing half of these individuals is sensible, because we have seen indicators suggesting that portfolio building tends to recommend a relatively diverse set of configurations (see Table 4.1). Yet, we still opt for an initial population partially consisting of random individuals to enforce diversity, because it is a crucial aspect for evolutionary search. The number of warm-starting individuals also affects the exploration-exploitation trade-off. The work introducing portfolio building provided results suggesting that 16 configurations may be a sensible value for warm-starting Bayesian optimization – an AutoML search procedure different from ours. Our setting differs not only on the AutoML search procedure, but also on the initialization because we include random configurations. Therefore, we cannot directly compare our larger number of warm-starting configurations (25) – and thus the exploration-exploitation trade-off – to the number of configurations used in the original work.

To evaluate warm-started AsyncEA we perform an analysis similar to the time-restricted setting. We perform a leave-one-dataset-out validation on OpenML-CC18: computing the method’s negative logistic loss on the left-out dataset using the metadata of all remaining datasets. However, now we evaluate the left-out dataset with a 3-fold cross-validation procedure as compared to hold-out. We include the meta-learning approach in the cross-validation setup – performing the meta-learner on all but the evaluation fold – but this is not required with the current dataset-agnostic meta-learner. Hence, we perform the 1-hour warm-started pipeline search multiple times per dataset – using different parts thereof – and average the results accordingly. We acknowledge that this evaluation setup is still not ideal; we only evaluate the method once on each part of the data, even though the search method is (partially) random in its initialization and search.

We compare the performance of warm-started AsyncEA with the non-warm-started 1-hour runs in the metadataset – i.e. the non-warm-started Random, Asynchronous Evolutionary and Asynchronous Successive Halving search methods. We rank the methods per dataset based on their cross-validation-based negative logistic loss scores. To test whether any of the optimization methods performs differently we employ the non-parametric Friedman test **<INCLUDE TEST RESULTS>**. Akin the time-restricted setting, we display the search method comparison in a critical differences (CD) diagram in Figure 4.6, employing a Wilcoxon signed-rank test considering a significance level $\alpha = 5\%$ and using a Bonferroni correction. Likewise, the methods’ dataset rank distributions are displayed in Figure 4.7.

test results

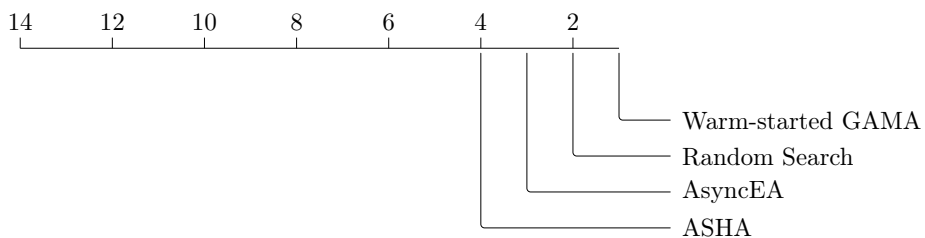


Figure 4.6: Critical differences (CD) diagram comparing warm-started and non-warm-started pipeline optimization approaches.

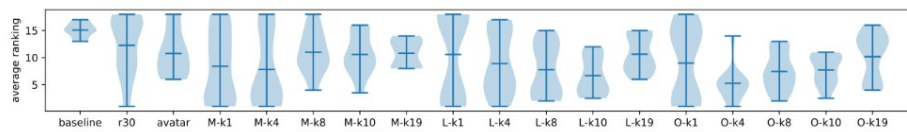


Fig. 1. The violin plots capturing the average rankings and ranking distribution for the configuration spaces on 11 datasets.

Figure 4.7: Example violin plot 2

Chapter 5

Conclusion and Discussion

Write your conclusions here.

Future work could focus on extending the toolbox with transferability metrics. And focus on incorporating tools for tasks other than classification, possibly extending to regression, or also unsupervised tasks. Hence, incorporating domain-similarity measures too in this toolbox. This would also allow to do source-selection for fine-tuning for example, it would allow for better NAS support.

Could also perform other analysis that are more focused on other parts, such as the metadatabase itself, and the effect of its construction on the outcomes.

Could include meta-learning approaches for two-layered AutoML tools like FLASH (fast bayesian optimization for data analytic pipelines).

(can quote Oracle AutoML paper) Generalizing knowledge transfer, deferring from implementation details, is an open research problem. It refers to the abstraction and generalization of a metadatabase, such that the prior created knowledge is transferable, regardless of the implementation details such as ML framework used, their versions, the version of the programming language, or the language itself. Also, the specific implementations of algorithms and so on.

Explore warm starting better, with different approaches to warm-start (like the amount of individuals etc). Potentially also with other meta-learning approaches and AutoML systems. Could also further explore the use of MLTA for other meta-learning applications such as runtime estimation or search space reduction.

Would have perhaps created a metadataset with varying amounts of optimization budget per dataset to not waste resources on overly exploring spaces, and to better generate prior experiences for the more difficult datasets.

Research whether we want to evaluate configurations before recommending them.

Currently the meta-learners are poorly parallelized, which could hurt performance when warm-starting search, because it takes up optimization time which does use the computational resources more effectively. Hence in future work it would be good to parallelize the meta-learner implementations.

Could in further developing the toolbox include (more) code tests, aiming for a code coverage of at least 90%.

Study warm-starting pipeline search for other type of automl framework (e.g. bayesian optimization). And could try to experiment with the exploration exploitation trade-off in warm-starting AutoML frameworks by both: the fraction of time given to the meta-learner and

really assess whether I have achieved the objectives I set out in CH1, perhaps sections

the amount of pipelines it recommends. Moreover could explore the impact of some diversity on the population, for instance to avoid having equal or very similar recommended pipelines. A possible approach for measuring diversity is by quantifying the (dis)similarity between pipelines by their characterizations, akin characterization-based dataset similarity measures.

Evaluation of meta-learners using dataset2vec may not be entirely valid, because dataset2vec is pre-trained with datasets that do re-occur in OpenML18CC.

The warm-starting potential score currently does not include the time it took, though this may effect the actual potential, we do not think that it would not distinguish between meta-learning approaches because the amount of meta-learner time is relatively small compared to the subsequent optimization time.

Should take into account RankML config char vs AP config char, but could better study whether the incorporation of hyperparameters helps in ranking pipelines, could relatively easily study this with RankML.

An interesting future research direction would be to explore how to warm-start, one such avenue could investigate learned defaults which are predicted using dataset characterizations, similarity, and possibly configuration characterizations and similarity thereover as well. It is much like "Meta-Learning for Symbolic Hyperparameter Defaults", but then for entire pipelines and possibly also for the meta-learner warm-starting something. We could thus have some sort of portfolio of good initializations for warm-starting, which could be learned and recommended using task-similarity. It is also much related to Part 2 of autosklearn 2.0:"to automatically select the best setting of the AutoML system itself for a given dataset"

Learned pipeline configurations, much like dataset2vec but for pipelines. But then of course having the ability of using some pre-trained model. So without having the high computational cost that factorization-based approaches have. That would be very nice.

In warm-starting now some recommended configurations may be incompatible with the dataset at hand, this could be solved with a pipeline checker such as AVATAR.

Study the robustness and scalability of the dataset characterization measures incorporated.

Referring to CH3 run time experiment: Also test robustness with multiple runs per dataset characterization method. Then the timings could also be aggregated over multiple runs which is better.

More research is perhaps needed to investigate whether Portfolio Building outperforms Dataset2Vec-based meta-learning approaches.

With the following in mind: "We suppose Portfolio Building's high configuration diversity is achieved by only considering the top performing configuration per dataset, thereby sampling configurations from 25 distinct datasets. The other meta-learning approaches also consider the configurations that did not achieve the best performance on the dataset, and are thus more likely to include configurations from a smaller number of datasets." It would be interesting to study the exploration exploitation trade off using the generalized Portfolio Building, allowing to use more than 1 configuration per dataset. We would expect that more configurations per dataset would reduce the diversity of solutions because many of the best solutions on each dataset tend to be similar.

Concerning the time-restricted analysis. Despite largely varying average rank statistics, we have found little statistically significant differences between meta-learning approaches. Therefore we can unfortunately not claim that certain meta-learning components outperform others. We postulate that with a higher statistical power (more measurements) we may be able to detect these differences. For the time-restricted analysis If I would have had more measurements - e.g. by cross-validation or including more datasets – then I would have

had more statistical power and may have detected performance differences between certain meta-learning components.

Its not really good to compare the 3-fold results against the 5-fold results, because the 5-fold results had more data to train on. Simply due to computational resource restriction it was not possible.

Support that we have achieved the research objective is CH4, but also the run time analysis on dataset characterization methods in CH3.

On a last note please observe that the influence of the amount of time and the number of recommended configurations on the trade-off may vary for each meta-learner and AutoML system. Therefore, investigating these effects is far from trivial and thus out-of-scope for the current study.

Something about also comparing the other meta-learning approaches in a warm-starting scenario, and why we have not done so.

The evaluation method in warm-starting is still not ideal, because you would want multiple runs.

Auto-sklearn actually shows that including the portfolio helps in warm-starting, but we cannot replicate this finding.

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Appendix A

Similarity Measure Usage in Meta-Learning Related Research Fields

Continual learning applications There is a mixed view on whether similar tasks help or hurt performance in continual learning. Some research shows improved performance using similar tasks, yet other argues for the use of dissimilar tasks [93].

One approach incorporates similarity information by extending continual learning approaches with a (1%) memory budget based on inter-task similarity [34]. The approach proves to be successful in mitigating catastrophic forgetting in low inter-task similarity settings. Another work proposes to incorporate a similarity detection function to analyze whether a specific task in the past is similar to the current task [130]. Then, the previous task’s knowledge is reused to slow down parameter expansion. Other research explored task-driven priors to choose modules for modular neural networks [128]. The method requires specification of the most similar task. Finding the most similar task is necessary because the search space grows exponentially. Once the most similar task is specified, the method evaluates the use of its modules for the current task to improve performance.

Multi-task learning A MTL study [99] shows competitive performance for pre-finetuning a smaller set of related tasks and large-scale multi-task learning. Moreover, the study finds that pre-finetuning on unrelated tasks diminishes performance. Thus, task relatedness, or similarity, is highly relevant for multi-task learning. Selecting an ideal subset of related tasks for multi-task pre-training is difficult [12], perhaps similarity measures help identify such groups of tasks. The incorporation of similarity in MTL has theoretical guarantees showing the benefit of explicit and implicit similarity knowledge [117]. Similarity measures may be suitable to provide the similarity for explicit-similarity MTL approaches. There are multiple works using similarity for weighting empirical loss [102, 91, 90]. Another study presents a multi-task learning approach relying on similarity using only features [74].

Neural Architecture Search (NAS) application Task-aware NAS [70] is a NAS framework employing an asymmetric similarity measure. The similarity measure is used to find the most similar tasks. Subsequently, the architecture search space is reduced by building upon the optimized architectures of, previously solved, similar tasks. Furthermore, there exists an approach to warm-start neural architecture search. More specifically, in [50] differentiable architecture search (DARTS) [76] is warm-started from similar tasks.

Curriculum learning applications Curriculum learning concerns ordering tasks, or data, in a meaningful order to improve performance, without increasing computational costs [119]. One study uses similarity information to effectively order the tasks by taking the shortest task through a task space [15]. Other work uses a similarity-based ordering to decompose a multi-task learning problem into a continual learning problem [103].

Synthetic data applications Another interesting line of research uses similarity information to generate synthetic data. One application is to choose an optimal data augmentation [11]. In data augmentation one applies transformations on the training data to achieve higher generalization performance. The research [11] shows a negative correlation between (augmented) dataset similarity and performance, it thus indicates that similarity measures may be used for ranking data augmentation procedures. Other research shows the use of representations learned by a similarity measure [86]. The approach uses similarity-measure-created representations to create synthetic data for pre-training [86]. The obtained performance is competitive with pre-training on Imagenet [30].

Drift detection and patch identification Concept drift approaches regularly utilize an active drift detection mechanism, informing when the data has changed substantially [56]. In this setup, the role of similarity measures is to help detect when drift occurs. To the best of our knowledge, there is little research establishing the use of similarity information in concept drift learning. We suspect this to be the case because concept drift learning is more concerned with similarities on a more fine-grained level, data, instead of datasets. We do believe that these methods can be adapted to incorporate similarity measures. There is one concept drift study relying on a similarity measure over chunks of data [22]. The study mentions the issue, at the time, of inadequate similarity estimation between chunks of data. Another study is connected to the previous application [26]. The study shows the applicability of similarity information for patch identification: determining whether two data patches are from the same initial dataset.

Transfer Learning The next few paragraphs are dedicated to a challenge in transfer learning, namely source selection for fine-tuning. Though transfer learning applies to a wide variety of models, it is best applied to neural networks because both their structure and parameters can be reused [126]. In transfer learning, we have source tasks and target tasks. Conventionally, source tasks have a remarkably higher cardinality than target tasks. Therefore, in transfer learning, an inductive bias is transferred from a source task to a target task. In the context of neural networks, the inductive bias usually consists of a model architecture and its trained parameters. Typically, there is more than one source task available, each with an associated pre-trained model. A common task in transfer learning is source selection for fine-tuning; a possible similarity measure downstream application.

Fine-tuning is a concept in transfer learning [124]. To understand fine-tuning it helps to think of neural networks consisting of two parts, a feature extractor and a top. The feature extractor extracts features from the raw input and feeds it to the top for classification or regression. There are different fine-tuning approaches, varying the amount the pre-trained model is changed [141]. We can copy the first n layers from the source to the target. The remaining layers of the target model are randomly initialized and trained on the target task. The amount of model change is controlled through n , yielding many fine-tuning options, potentially with different rates of change per layer [111]. Simplifying this choice, typically

one of two approaches is used. Either fine-tune the feature extractor, backpropagating the target task’s errors, or freeze the feature extractor, leaving its values unchanged. This choice depends on the similarity between the source and target task. A popular approach is to freeze the feature extraction layers and fine-tune the classification or regression top. Fine-tuning is powerful because it allows for training complex neural networks, with relatively little data, in turn achieving otherwise unachievable high performance [81, 124].

In source selection for fine-tuning, the goal is to select the source task, or model, which after fine-tuning yields the best performance on the target task. An additional goal is to select the source task efficiently, on that account, it is prohibited to continue training the source models on the target task. Similarity measures are well-suited to the source selection task because more similar source and target tasks achieve higher transferability [141]. Even more so, in transfer learning, knowledge transfer yields degrading performance if the source and target are too dissimilar [131, 144]. An interesting extension of source selection is multi-source feature fusion, which has been proven to benefit from similarity information [120]. In multi-source feature fusion, multiple source models are transferred to the target task by merging their inferred features to obtain a fused representation.

Similarity as weight Active testing [72] is an approach to classification algorithm selection employing dataset similarity. Active testing is initialized with a high-performing default and proceeds tournament style. In rounds, it selects and evaluates the configurations most promising to beat the current best solution. In identifying the most promising competitor a similarity measure is used in weighting. The active testing framework for algorithm selection can likely benefit from other, more recent, similarity measures.

Apart from transferring configurations (see Section 2.2.2.1), we can also transfer surrogate models. A surrogate model models the evaluations of the configurations. A straightforward approach to transfer a surrogate model is to transfer the one from the most similar task. A general and powerful approach to transferring surrogate models is to transfer multiple of them [40], potentially weighted by a similarity value [135]. In [135] the authors propose a similarity measure based on landmark vectors, but the overall approach is flexible enough to function with any similarity measure.

Appendix B

Supplemental Material Metadataset

Table B.1: The number of configuration evaluations for each dataset in OpenML18CC.

OpenML Dataset Name	OpenML ID	Number of Evaluations
ilpd	1480	76945
blood-transfusion-service-center	1464	70066
balance-scale	11	56495
climate-model-simulation-crashes	1467	49823
breast-w	15	49360
diabetes	37	48927
banknote-authentication	1462	48181
tic-tac-toe	50	44684
kc2	1063	42227
dresses-sales	23381	37862
analcatdata_dmft	469	33786
analcatdata_authorship	458	24003
pc3	1050	23880
credit-approval	29	23492
cmc	23	18620
qsar-biodeg	1494	18509
wilt	40983	17997
vehicle	54	17030
car	991	16233
kc1	1067	16084
pc4	1049	15973
mfeat-morphological	18	15248

credit-g	31	14719
MiceProtein	40966	14189
cylinder-bands	6332	14177
pc1	1068	13991
vowel	307	13275
eucalyptus	188	12151
wdbc	1510	11939
spambase	44	11912
phoneme	1489	10011
sick	38	8252
jm1	1053	7515
ozone-level-8hr	1487	7409
kr-vs-kp	3	7112
steel-plates-fault	1504	6641
segment	36	5159
churn	40701	4456
mfeat-fourier	14	4381
mfeat-karhunen	16	4111
mfeat-zernike	22	3890
PhishingWebsites	4534	3874
jungle_chess_2pcs_raw_endgame_complete	41027	3593
electricity	151	3280
wall-robot-navigation	1497	3188
GesturePhaseSegmentationProcessed	4538	2487
satimage	182	2395
optdigits	28	2381
pendigits	32	2032
dna	40670	1872
bank-marketing	1461	1739
texture	40499	1725
first-order-theorem-proving	1475	1630
mfeat-pixel	20	1536
mfeat-factors	12	1475
splice	46	1465
semeion	1501	1308

letter	6	1107
Internet-Advertisements	40978	952
adult	179	925
numerai28.6	23517	905
madelon	1485	898
cnae-9	1468	892
nomao	1486	639
connect-4	1591	534
isolet	300	503
Bioresponse	4134	439
har	1478	411
mnist_784	554	233
Fashion-MNIST	40996	207
Devnagari-Script	40923	114
CIFAR_10	40927	21

Appendix C

Common Meta-Feature Collections

Table C.1: Table depicting two common and well-performing collections of meta-Features for supervised classification tasks, namely the meta-features and summarization functions by Feurer et al. [43] and those by Wistuba et al. [134]. Note that meta-features are grouped by their categorization and that the Wistuba meta-features are a subset of the Feurer meta-features.

Meta-Feature Group	Feurer Meta-Features	Wistuba Meta-Features
Simple	Number of instances	Number of instances
	Log number of instances	Log number of instances
	Number of classes	Number of classes
	Number of features	Number of features
	Log number of features	Log number of features
	Number of instances with missing values	
	Percentage of instances with missing values	
	Number of features with missing values	
	Percentage of features with missing values	
	Number of missing values	
	Percentage of missing values	
	Number of numeric features	
	Number of categorical features	
	Ratio numerical to categorical	
	Ratio categorical to numerical	
	Dataset dimensionality	Dataset dimensionality
	Log dataset dimensionality	Log dataset dimensionality
	Inverse dataset dimensionality	Inverse dataset dimensionality
	Log inverse dataset dimensionality	Log inverse dataset dimensionality
	Class probability min	Class probability min
	Class probability max	Class probability max
	Class probability mean	Class probability mean
	Class probability std	Class probability std

Information-theoretic	Class entropy	Class entropy
Statistical	Min number of categorical values	
	Max number of categorical values	
	Mean number of categorical values	
	Std number of categorical values	
	Total number of categorical values	
	Kurtosis min	Kurtosis min
	Kurtosis max	Kurtosis max
	Kurtosis mean	Kurtosis mean
	Kurtosis std	Kurtosis std
	Skewness min	Skewness min
	Skewness max	Skewness max
	Skewness mean	Skewness mean
	Skewness std	Skewness std
PCA	PCA 95%	
	PCA skewness first principal component	
	PCA kurtosis first principal component	
Landmarking	One Nearest Neighbor	
	Linear Discriminant Analysis	
	Naive Bayes	
	Decision Tree	
	Decision Node Learner	
	Random Node Learner	

Appendix D

OpenML-CC18 Metadataset Subset

Table D.1: The OpenML18CC subset. Datasets were selected by hand to not overlap with the datasets that were used for training Dataset2Vec [58].

OpenML Dataset Name	OpenML ID
climate-model-simulation-crashes	1467
banknote-authentication	1462
kc2	1063
dresses-sales	23381
analcatdata_dmft	469
analcatdata_authorship	458
pc3	1050
qsar-biodeg	1494
wilt	40983
kc1	1067
pc4	1049
mfeat-morphological	18
MiceProtein	40966
pc1	1068
eucalyptus	188
phoneme	1489
jm1	1053
kr-vs-kp	3
churn	40701
mfeat-fourier	14
mfeat-karhunen	16
mfeat-zernike	22
PhishingWebsites	4534
jungle_chess_2pcs_raw_endgame_complete	41027
electricity	151

GesturePhaseSegmentationProcessed	4538
first-order-theorem-proving	1475
mfeat-pixel	20
mfeat-factors	12
Internet-Advertisements	40978
numerai28.6	23517
madelon	1485
cnae-9	1468
nomao	1486
isolet	300
Bioresponse	4134
har	1478
mnist_784	554
Fashion-MNIST	40996
Devnagari-Script	40923
CIFAR_10	40927

Appendix E

Meta-Learning Approaches' Regret Formalization

Let \mathcal{M} be the set of all meta-learning approaches and \mathcal{D}_E the set of all evaluation datasets. Let a meta-learning approach $m_i \in \mathcal{M}$ be tasked with composing a warm-starting vector $\mathbf{w}_{i,j} = [(g, \mathbf{A}, \boldsymbol{\lambda})_k, (g, \mathbf{A}, \boldsymbol{\lambda})_{k+1}, \dots, (g, \mathbf{A}, \boldsymbol{\lambda})_{N_C}]$ consisting of N_C pipeline configurations for dataset $d_j \in \mathcal{D}_E$. Associated to the warm-starting vector $\mathbf{w}_{i,j}$ is the evaluation vector $\mathbf{e}_{i,j} \in \mathbb{R}^{N_C}$ consisting of the evaluation scores for executing the pipeline configurations in $\mathbf{w}_{i,j}$ on dataset d_j , thus $\mathbf{e}_{i,j,k}$ denotes the evaluation score for meta-learning approach m_i using its k^{th} recommended configuration on dataset d_j . For normalization we create reference evaluation scores $t_j, b_j \in \mathbb{R}$ for each dataset $d_j \in \mathcal{D}_E$ composed of the top and bottom pipeline configuration evaluation scores respectively, thus

$$t_j = \max_{i,k}(\mathbf{e}_{i,j,k}), \quad b_j = \min_{i,k}(\mathbf{e}_{i,j,k}) \quad (\text{E.1})$$

If meta-learning approach m_i fails to provide a configuration recommendation $(g, \mathbf{A}, \boldsymbol{\lambda})_k$ in $\mathbf{w}_{i,j}$ its entry $\mathbf{w}_{i,j,k}$ will be empty, and its corresponding evaluation entry $\mathbf{e}_{i,j,k}$ is set to b_j to yield $\frac{|t_j - \mathbf{e}_{i,j,k}|}{|t_j - b_j|} = \frac{|t_j - b_j|}{|t_j - b_j|} = 1$. Assuming the evaluations in \mathbf{e} are from a scoring function (higher is better), we can assign each meta-learning approach m_i its average normalized regret:

$$r_i = \left(\frac{1}{|\mathcal{D}_E| \cdot N_C} \sum_{j=1}^{|\mathcal{D}_E|} \sum_{k=1}^{N_C} \frac{|t_j - \mathbf{e}_{i,j,k}|}{|t_j - b_j|} \right) \in [0, 1] \quad (\text{E.2})$$

which can be interpreted as the average normalized Manhattan distance between the worst possible and the recommended configuration scores, averaged over all datasets. Or in other words it is the normalized regret, averaged over datasets and recommended configurations.