

Declarative Metadata Management: A Missing Piece in End-To-End Machine Learning

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ABSTRACT

We argue for the necessity of managing the metadata and lineage of common artifacts in machine learning (ML). We discuss a recently presented lightweight system built for this task, which accelerates users in their ML workflows, and provides a basis for comparability and repeatability of ML experiments. This system tracks the lineage of produced artifacts in ML workloads and automatically extracts metadata such as hyperparameters of models, schemas of datasets and layouts of deep neural networks. It provides a general declarative representation of common ML artifacts, is integrated with popular frameworks such as MXNet, SparkML and scikit-learn, and meets the demands of various production use cases at Amazon.

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1 INTRODUCTION

When developing and productionizing ML models, a major proportion of the time is spent on conducting model selection experiments which consist of training and tuning models and their corresponding features [2, 3, 13, 19, 22, 27]. Typically, data scientists conduct this experimentation in an ad-hoc style without a standardized way of storing and managing the resulting experimentation data and artifacts. As a consequence, the results of these experiments are often not comparable, as there is no standard way to determine whether two models had been trained on the same input data, for example. Even more, it is tedious and time-consuming to repeat successful experiments later in time, and it is hard to get an overall picture of the progress made in ML tasks towards a specific goal, especially in larger teams. Simply storing the artifacts (datasets, models, feature sets, predictions) produced during experimentation in a central place is unfortunately insufficient to mitigate this situation. Achieving repeatability and comparability of ML experiments forces one to understand the metadata and, most importantly, the lineage of artifacts produced in ML workloads [13]. For example, in order to re-use a persisted model, it is not sufficient to restore its contents byte by byte; new input data must also be transformed

into a feature representation that the model can understand, so information on these transforms must also be persisted. As another example, in order to reliably compare two experiments, we must ensure that they have been trained and evaluated using the same training and test data respectively, and that their performance was measured by the same metrics.

To address the aforementioned issues and assist data scientists in their daily tasks, we proposed a lightweight system for handling the metadata of ML experiments [20]. This system allows for managing the metadata (e.g., *Who created the model at what time? Which hyperparameters were used? What feature transformations have been applied?*) and lineage (e.g., *Which dataset was the model derived from? Which dataset was used for computing the evaluation data?*) of produced artifacts, and provides an entry point for querying the persisted metadata. Data scientists can leverage this service to enable a variety of previously hard-to-achieve functionality, such as regular automated comparisons of models in development to older models (similar to regression tests for software). Additionally, the proposed service helps data scientists to easily ad-hoc test their models in development and provides a starting point for quantifying the accuracy improvements that teams achieve over time towards a specific ML goal, e.g., by storing and analyzing the evaluation results of their models and showing them via a leaderboard. In order to ease the adoption of our metadata tracking system, we explore techniques to automatically extract experimentation metadata from common abstractions used in ML pipelines, such as ‘data frames’ which hold denormalized relational data, and ML pipelines which comprise a way to define complex feature transformation chains composed of individual operators. For applications built on top of these abstractions, metadata tracking should not require more effort than exposing a few data structures to our tracking code.

In the following, we summarize the design decisions for our system (Section 2), list related work (Section 3) and discuss future research directions for ML metadata management (Section 4).

2 SYSTEM DESIGN

Data model. The major challenge in designing a data model for experimentation metadata is the *trade-off between generality and interpretability of the schema*. The most general solution would be to simply store all data as bytes associated with arbitrary key-value tags. Such metadata however would be very hard to automatically interpret and analyze later, as no semantics are enforced. A too narrow schema on the other hand might hinder adoption of our service, as it does not allow scientists to incorporate experimentation data from a variety of use cases. We propose a middle ground with a schema¹ that strictly enforces the storage of lineage information (e.g., which dataset was used to train a model) and ML-specific

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¹available at <https://github.com/awsml/ml-experiments-schema>

attributes (e.g., hyperparameters of a model), but still provides flexibility to its users by supporting arbitrary application-specific annotations. The most important principle we embrace is *declarativity*: we store metadata of the artifacts but not code that produces it, and only store pointers to the actual input data or serialized parameters. This enforces a strict decoupling, and enables querying and analysis of the metadata and lineage. The second important principle for our system is *immutability*: metadata entries are only written once, ruling out a variety of potential consistency problems. **Architecture.** Our system employs a three-layered architecture: On the lowest layer, a document database stores the actual experimentation data. In the next layer, this centralized data store is exposed to the outside world via a REST API, for which we provide so-called low-level clients for the JVM and Python, which allow users to explicitly store metadata for particular artifacts and query the existing data. The uppermost layer is formed by what we call high-level clients that are geared towards popular ML libraries such as *SparkML* [14], *scikit-learn* [17], *MXNet* [4].

Automated Metadata Extraction. Our high-level clients enable automated metadata extraction from internal data structures of popular ML frameworks. ML workloads in *SparkML* for example are comprised of pipeline stages which operate on *DataFrames*, a relational abstraction for a partitioned table with a well-defined schema. The architecture of *SparkML* pipelines allows us to automatically track all the schema transformations (e.g. reading, adding and removing columns) each pipeline operator conducts, as well as the parameterization of the operators. We create and store directed acyclic graph representations of *Spark* pipelines (where edges denote pipeline operators and vertices correspond to dataframe columns) by extracting the schema of the input data frame and replaying the schema changes the pipeline conducts. Frameworks for deep neural networks offer their users a very fine-grained abstraction to declaratively define their models by combining mathematical operators (e.g., linear layers, convolutions, activation functions) into the layout of the network to learn. Our high-level client for *MXNet* [4] for example extracts and stores the resulting computational graph, together with the parameterization and dimensionality of the contained operators, and corresponding hyperparameters such as optimizer settings.

3 RELATED WORK

In order to foster collaboration between scientists, platforms such as *OpenML* [25] and the *W3C ML Schema* initiative [10] allow researchers to share descriptions and evaluation results of their ML experiments. Managing and efficiently executing model selection workloads has been identified as an upcoming challenge [11, 13, 21] in the data management community. The *ModelDB* [26] project puts a specific focus on organizing models, and comes very close to our system design-wise with the difference that we support more general classes of models and apply more detailed tracking. Other classes of systems specialize on deep learning [16], aim at efficiently serving the resulting models for prediction [5, 6] or concentrate on tracking and indexing provenance information [7, 15, 18]. Modeling ML workloads via pipelines (which are typically inspired by the ‘estimator/transformer’ abstraction in *scikit-learn* [17]) and efficiently executing such pipelines at scale has become an active area of

research. Established work describes production systems and platforms [2, 3], investigates software engineering aspects [22, 24] and pipeline abstractions for machine learning workloads [1], often on top of the massively parallel dataflow system *Apache Spark* [14, 23].

4 OUTLOOK

We see a huge potential in enabling declarative management of ML metadata: data scientists are provided with infrastructure that allows them to accelerate their experimentation via dashboards and leaderboards that list experiments, email notifications which summarize experimentation progress, and automated regression tests for the prediction quality of ML models during development, which compare the results on hold out data with historical prediction results, e.g. upon every commit to the codebase. In the remainder, we elaborate on research directions and upcoming challenges for ML metadata management systems:

Replicable Model Training and Deployment. Systems like ours allow companies to accelerate their experimentation and innovation cycle, and have the potential to form a corner stone of replicable ML model training, which will become more important in the light of upcoming legal requirements for the real-world usage of machine learning. In order to enable such a replicable model training, it is not sufficient to be able to access the metadata of the ML workload, additionally the training source code (e.g., via the corresponding git commit) as well as the computational environment (e.g., via a docker image) need be tracked and stored. Ideally such a system would not only automate replicable model training but also integrate the resulting models with model serving systems for easy deployment.

Further Automation of Metadata Tracking. Although we provide elaborate extraction functionality, we currently still rely on users understanding our complex schema and correctly integrating their code with our API. We aim to increase the automation of our extraction code and to decrease the amount of additional code and effort required to enable the metadata tracking in a workload. One direction to investigate is the instrumentation of notebooks like *Jupyter* [12], where we would ideally assist the user interactively in tracking ML metadata during explorative model tuning efforts [18]. Another orthogonal direction would be to extract ML metadata posthoc from logfiles produced by model training systems.

Meta Learning. Our long term research goals include enabling meta learning [8] on top of our experiment repository, e.g. to recommend features, algorithms or hyperparameter settings for new datasets. This would require us to implement the automated computation of *metafeatures* [9] for contained datasets, as well as similarity queries allowing users to find the most similar datasets for new data, based on these metafeatures. It would furthermore be beneficial to additionally leverage data from open repositories such as *OpenML* [25] for this task.

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