Smart Model Elimination for Efficient Automated Machine Learning in Healthcare

Eric Su Zhang¹

St. Mark's School of Texas 10600 Preston Rd, Dallas, TX 75230 26zhange@smtexas.org

Benjamin Joseph Michael Standefer²

St. Mark's School of Texas 10600 Preston Rd, Dallas, TX 75230 26standeferb@smtexas.org

Abstract

Automated Machine Learning (AutoML) has emerged as a popular field of research. We present a literature review of existing AutoML papers and conduct a survey on five popular AutoML frameworks. Typically, these frameworks engage in model selection by requiring every model to be run and assessed, a process both time-intensive and computationally expensive. In response, we propose a novel framework, Smart Model Elimination Machine Learning (SMEML), that strategically eliminates models that are unlikely to yield high accuracy. SMEML demonstrates the ability to achieve comparable accuracy to a traditional brute-force approach while significantly reducing the time required. We also believe this innovation is particularly beneficial to machine learning in healthcare, where efficient and accurate disease prediction is crucial.

1 Introduction

Machine learning (ML) is a type of artificial intelligence (AI) that uses searching and trend-determining to make predictions. ML algorithms are trained and tested on a given dataset to become "smarter". Once they've learned the patterns typical of that data, they can make predictions based on similar data. These algorithms also require additional processes, however, like hyperparamater tuning, and can be modified in conjunction with one another, like in ensemble learning. This makes the process complex and time-inefficient. To do this on a larger scale and at greater speeds, recent research and effort have been put into developing AutoML frameworks [1].

One sector of the global industry that has the most potential for beneficial integration of AutoML frameworks is healthcare, particularly in developing countries. These countries have severe shortages of healthcare workers and limited tools for diagnosis. For example, Africa has 2.3 healthcare workers per 1000 individuals, while the Americas have 24.8 healthcare workers per 1000 [2]. The World Health Organization (WHO) emphasized that this deficit is growing every year and will likely reach 18 million personnel by 2030 [3]. Medical AIs typically automate repetitive tasks, making time consumption a primary concern [4]. This coupled with a lack of adequate resources makes designing faster diagnostic systems for developing countries' medical sectors a pivotal issue.

Most AutoML frameworks implement some form of model selection where a pool of models are filtered until a final model is selected. However, these frameworks require that every model be run [1]. In theory, this means that much energy is spent training models that are unlikely to be selected. We propose SMEML, a novel model selection algorithm that automatically eliminates models that it believes will not be performant.

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Table 1: Survey of Existing AutoML Frameworks

Framework	EOU	M	FE	TM	HPT	Inter	Custom
H2O AutoML [5]	Easy	12	Basic	Yes	Auto	SHAP	High
TPOT [6]	Hard	10	Basic	No	Genetic	POJO	Moderate
MLJAR [7,8]	Easy	12	Advanced	Yes	Auto	Basic	High
FLAML [9]	Easy	9	Limited	Yes	Auto	SHAP	Low-Moderate
LightAutoML [10]	Easy	11	Advanced	Yes	Auto	Basic	Moderate

Abbreviations: EOU: Ease of Use, M: # of models evaluated by framework, FE: Feature Engineering, TM: Time Management, HPT: Hyper-Parameter Tuning, Inter: Interface, Custom: Customization

1.1 Survey

We systematically review and compare five different AutoML frameworks, comparing every feature relevant to medical implementation in Table 1. Of these five, we wanted to take a closer look at four: FLAML, H2O, MLJAR, and TPOT.

We can observe pros and cons in the four models' theoretical applications. FLAML uses blend search hyper-parameter optimization and focuses on lightweight models, making it quick and applicable to repetitive tasks. It supports user-specified imputation techniques for missing values and has an intuitive API, making it moderately user-friendly for healthcare workers [9]. H2O has a wider variety of supported algorithms that may be appropriate for medical diagnosis, making it slower than FLAML on average. In one trial, more than half of FLAML's performances in one minute were better than or equal to H2O's performances in one hour [9]. H2O has automatic and multi-faceted methods of dealing with missing values and flexible interfaces in R and Python, making it intuitive for users [5]. MLJAR uses advanced algorithms such as light gradient boosting and neural networks. It automatically deals with missing values and is known for its automated user interface [7]. TPOT uses genetic programming. It builds pipelines over multiple generations, which can be time consuming. It has built-in pre-processing for missing values, but the genetic programming requires fine-tuning from users [6].

Through this survey, we uncovered a key continuity in these frameworks' design: frameworks must run all of their models to maximize accuracy at the expense of efficiency. Given the specific nature of certain tabular data, there are some models that are less likely to perform well, thus running and training these models is wasteful. Cutting these models completely, however, is not optimal, as they are the best option in a minority of cases. We hypothesized that using a combination of multi-layer machine learning implementation and meta-feature extraction would yield an equally versatile system that minimizes time spent on training without sacrificing accuracy.

Methodology

When considering a specific dataset, the distribution, relationship between datapoints, and various other attributes can give us clues as to which models are likely to perform best. For example, linear models tend to work better on datasets where there is a linear relationship between features [11]. Accordingly, we can develop a boosting model to predict the best models based on extracted attributes from a dataset. These extracted attributes are shown in Table 2. Each attribute attempts to capture a specific type of relationship or distribution within the data.

We considered 28 different models, including all supervised and semi-supervised sklearn [12] classification models and common boosting models such as XGBoost [13], LightGBM [14], and Catboost [15].

To implement the model elimination process, we train a multilabel-regression boosting model called the SME Model-Ranker on 268 kaggle datasets. We choose medical datasets and focus our approach on binary classification. We extract the attributes in Table 2 from each dataset: which are the inputs for the boosting model. We then run a "dumb", brute-force framework that trains every model every each dataset. The accuracy of every model is ranked, and every model is treated as a label with its value being its rank. We train the SME to predict the rank of each model, given the extracted attributes. SME produces a median spearman correlation of 0.7301 between the predicted rank and the actual rank and a median p-value of 1.032e-05. Furthermore, we guarantee that we can predict a top two model within the predicted top eight 90.12% of the time.

Table 2: Attributes extracted from a dataset

Attribute	Formula	Purpose	Weight
# of Rows	$ \mathbf{A} _r$	DS	.0422
# of Columns	$ \mathbf{A} _c$	DS	.0415
Target Distribution	$\frac{\max(count_1, count_0)}{ \mathbf{A} _r}$	DT	.0489
Numerical Columns	$\frac{ \mathbf{A}_{num} }{ \mathbf{A} _c}$	DT	.0428
Binary Categorical Columns	$rac{ \mathbf{A}_{bin} }{ \mathbf{A} _c}$	DT	.0117
Mean # of Distinct Values	μ {# of distinct values in column}	DT	.0076
Mean IQR	μ {IQR of column}	DD	.0594
STD of IQR	σ {IQR of column}	DD	.0700
Mean Q1	μ {Q1 of column}	DD	.0789
STD of Q1	σ {Q1 of column}	DD	.0613
Mean Q3	μ {Q3 of column}	DD	.0812
STD of Q3	σ {Q3 of column}	DD	.1103
Mean Percent of Z-Score Outliers	$\mu \left\{ \frac{\text{\# of Z-Score Outliers}}{ \mathbf{A} _r} \right\}$	ODP	.0558
STD of Percent of Z-Score Outliers	$\sigma \left\{ \frac{\text{# of Z-Score Outliers}}{ \mathbf{A} _r} \right\}$	ODP	.0798
Mean Correlation	μ {Correlation Matrix}	DL	.0559
STD of Correlation	σ {Correlation Matrix}	DL	.1526

Abbreviations: DS: Data Size, DT: Data Type, DD: Data Distribution, ODP: Outlier Data Points, DL: Data Linearity, Weight: Variable Importance

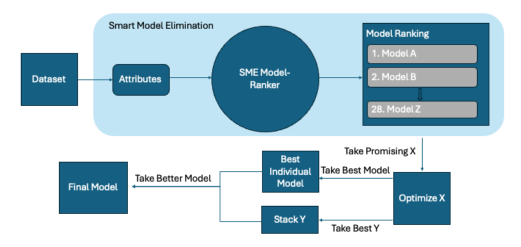


Figure 1: Illustration of the SMEML Process

When a user inputs a dataset into SMEML, it first extracts the attributes from the dataset and then feeds them into the SME to get the predicted rank of each model. Since we have determined that we can predict a top two model within the predicted top eight quite confidently, we can eliminate the bottom 20 models from the list. We run Bayseian optimization using scikit-optimize [16] on the top eight models to get the best hyperparamters for each model. We select the top models that have significantly better performance than other models. These models are stacked together and compared to the best individual model. The final model is the better one out of these two. This process is shown in Figure 1.

Table 3: Comparing experiment results for SMEML and dumb mode

Dataset	Accuracy		Time to Train (sec)		Size (RxC)
	SMEML	Dumb	SMEML	Dumb	
Lung Cancer Prediction [17]	.9839	.9839	143.07	317.04	309x16
Chronic Kidney Disease Dataset [18]	1.0	1.0	419.64	788.46	400x26
Pima Indians Diabetes Dataset [19]	.7857	.8052	161.99	374.01	768x8
Brain Stroke Prediction Dataset [20]	.9458	.9458	114.16	370.95	4892x11
Stroke Prediction Dataset [21]	.9393	.9393	172.67	385.23	5110x12
Eye State Classification [22]	.9536	.9549	312.57	696.51	14980x15

3 Evaluation

3.1 Experiment Design

To evaluate our framework, we choose 5 different datasets of diverse size. We ran SMEML on each dataset for 20 iterations of tuning. We compared this against a "dumb" mode which simply runs every model on the dataset. We used the accuracy and time to train as our metrics. All experiments are run on an Ubuntu server with Intel Xeon E5520 @ 2.27GHz and 24GB of RAM.

3.2 Experiment Results

Several observations were made from running these datasets. First, accuracy did not change for 4 of the 6 benchmarks. The other two benchmarks on eye states and diabetes prediction represented a loss of accuracy for SMEML by 0.13% and 1.95%, respectively. Training time, however, favored SMEML in every trial. SMEML outperformed the dummy framework by 173.96 seconds in its worst performance on the lung cancer trial and by 383.94 seconds in its best performance on the eye state dataset. Relative to the dummy's performance, a percent decrease in time to run for SMEML of at least 46.78% could be seen, with a decrease of up to 69.22% observerd as well. SMEML's worst performance by percent decrease was also the performance in which it returned perfect accuracy.

4 Discussion

4.1 Summary

SMEML consistently achieves comparable accuracy to its dumb counterpart while significantly reducing the time required to train the model. Notably, SMEML's performance is not limited by the size of the dataset, achieving its best runtimes–first, third, and fourth fastest overall—on 'large' datasets (those exceeding 1000 rows). This efficiency is important in third world medicine where there is a constant influx of new patient data that must be processed quickly. A practical demonstration of this software's real-world application can be found on our SmartDx website [23]. Source code can be found at [24].

4.2 Limitations

Our implementation of SMEML has limitations. These include a small training set due to lack of available datasets, finite computing power, time constraints, and a lack of access to paticular scientific literature. Our framework only improves on the model selection process and does not address the various other aspects of AutoML. Therefore, SMEML is not comparable to other full-featured AutoML frameworks.

4.3 Future Work

Future work will include increasing the size of the training set, implementing a fully fledged AutoML framework, and conducting a detailed experiment comparing SMEML to other popular AutoML frameworks. Furthermore, we will examine other types of machine learning such as regression and multiclass-classification.

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