Towards Comparable Active Learning

David S. Hippocampus*

Department of Computer Science Cranberry-Lemon University Pittsburgh, PA 15213 hippo@cs.cranberry-lemon.edu

Abstract

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5 1 Introduction

Among others, the authors of [14] have pointed out severe inconsistencies in results of AL papers in recent years. In their supplementary materials they conducted a meta analysis of reported results of several different AL algorithms and found that all considered algorithms only provided significant lifts in their own original papers, while all following literature reported performances no better that uncertainty sampling, or in some cases no better than random sampling for the same algorithm. The result of these inconsistencies is a chaotic landscape of AL algorithms where every paper claims to archive state-of-the-art results by significantly outperforming everyone else, while the vast majority of results proves to be non-reproducible.

14 1.1 Contributions

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- 1. Evaluation of Active Learning algorithms on datasets from 4 different domains, including synthetic data that highlights principled shortcomings of existing approaches.
- 2. Novel experimental protocol for seeding the experiment with 3 different seeds to allow full control and reproducibility and analysis of how many restarts are required to converge to the true median performance reliably.
- Simple algorithm for an Oracle-Curve that can be constructed greedily and does not rely on search.

22 **Related Work**

- Version: Braindump
- 24 Many different algorithms have been proposed for active learning. In this work we focus on those
- 25 approaches that have shown consistent results over the years as well as newer approaches that have
- 26 demonstrated significant lifts in their initial experiments. AL algorithms can be categorized into

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clude CoreSet [12] and TypiClust [4], which use clustering techniques to partition the data and then 28 29 sample their unlabeled points based on the clusters. Uncertainty-based approaches include classic uncertainty sampling (based on Shannon-Entropy and the margin-score), BALD [6] and BADGE 30 [1], which use metrics to measure the classifiers state. 31 32 Some previous work also aimed to provide a benchmark suite for active learning: The authors of [2] and [8] both focus on active learning in the image domain. While [2] discuss a new metric to 33 measure AL performance, which they call "Label Efficiency" and provide experiments on many 34 common configurations for data preparation, model training and other hyperparameters, [8] focuses 35 on combined approaches of AL and semi-supervised learning to aid model training. The authors of 36 [5] study models that are learned with AL techniques in the image and text domain. They test for 37 several different properties of the models including robustness, response to compression techniques 38 and final performance. 39

two classes: Geometric approaches and uncertainty-based approaches. Geometric approaches in-

Overview

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- Table 1 shows a feature comparison between our proposed benchmark and several existing bench-
- marks in the literature, as well as methodological AL papers with an extensive experiments section.
- TODO We include in this table any methodological paper that experiments on at least two domains. TODO Define AL scenarios (really hard)

Paper	# Datasets	Domains	Scenarios	Oracle	RL Alg.
Beck et al. [2]	4	1	3	-	-
Hu et al. [5]	5	2	1	-	-
Li et al. [8]	5	1	1	-	-
Zhou et al. [14]	3	2	1	\checkmark	-
Ours	5	2	2	\checkmark	-

Table 1: Comparison of our benchmark with the existing literature

3.1 Problem Description

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- Given two spaces $\mathcal{X} := \mathcal{R}^M$ and $\mathcal{Y} := \mathcal{R}^C$, a sample $\mathcal{D}_1, \dots, \mathcal{D}_N \subseteq (\mathcal{X} \times \mathcal{Y})^*$ of sequences of 47
- pairs (x, y) from an unknown distribution p called datasets and a number $B \in \mathcal{N}$ with $B < |\mathcal{D}|$.
- Given two functions $\ell: \mathcal{Y} \times \mathcal{Y} \to \mathcal{R}$ called loss, and $A: (\mathcal{X} \times \mathcal{Y})^* \times \mathcal{X}^* \to \mathcal{Y}^{\mathcal{X}}$ called learning
- algorithm, find a function

$$a:(\mathcal{X}\times\mathcal{Y})^*\times\mathcal{X}^* o\{0,1\}^*$$
 (equivariant in the second argument)

called acquisition function, s.t. the expected loss of a model learned on all predictors plus B acquired targets is minimal:

$$\begin{split} \mathbb{E}_{\mathcal{D} \text{train}, \mathcal{D} \text{test} \sim p} & \operatorname{avg}_{(x,y) \in \mathcal{D} \text{test}} \ell(y, \hat{y}(x)) \\ & \text{with } \hat{y} := & A((\mathcal{D} \text{train}_{n_1}, \dots, \mathcal{D} \text{train}_{n_B}), \mathcal{D} \text{train}|_{\mathcal{X}}) \\ & n_b := & \operatorname{index}(a((\mathcal{D} \text{train}_{n_1}, \dots, \mathcal{D} \text{train}_{n_{b-1}}), \mathcal{D} \text{train}|_{\mathcal{X}})), \quad b \in 1:B \end{split}$$

- (We would need to switch from lowest expected loss to highest AUC) 53
- Since combinatorial problem of finding the optimal subset \mathcal{D}_{train} is computational not feasible, we 54 55
- allow sequential construction of \mathcal{D}_{train} .
- To construct the active learning setting, we suppress the labels \mathcal{Y} of \mathcal{D}_{train} to form the unlabeled 56
- pool $\mathcal{U}:=\mathcal{X}^*$ and form and initial labeled pool by uniformly sampling k number of instances per 57
- class from \mathcal{U} and recovering their label $\mathcal{L} := (\mathcal{X}, \mathcal{Y})^{k*C}$. 58

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Following [14], the quality of an active learning algorithm is evaluated by an "anytime" protocol that incorporates classification performance at every iteration, not just the final performance after the budget is exhausted. We employ the normalized area under the accuracy curve (AUC):

$$\operatorname{auc}(\mathcal{U}, \mathcal{L}, \hat{y}, B) := \frac{1}{B} \sum_{i=1}^{B} \operatorname{Acc}(y_{test}, \hat{y}_i(x_{test}))$$
(1)

Where \hat{y}_i is the retrained classification model after the i-th instance was selected.

64 Framing AL as RL

We define the active learning process as an adapted reinforcement learning loop $(S, A, \tau, \Omega, \omega)$ where an environment iteratively will expose a state $s \in S$ to an agent Ω , which will choose actions $a \in A$. For each iteration i the environment samples a subset of size τ of unlabeled instances $u^{(i)} \sim \mathcal{U}^{(i)}$, constructs the state $s^{(i)} := \omega(u^{(i)})$ and presents it to the agent to select an action $a^{(i)} := \Omega(s^{(i)})$. The action $a^{(i)}$ is the index of the selected instance in $u^{(i)}$ out of all possible indices $A := [1 \dots \tau]$. This process is repeated B times $i := [1 \dots B]$.

Algorithm 1 Active LearningRequire: \mathcal{U} \triangleright Unlabeled PoolRequire: τ \triangleright Unlabeled Sample SizeRequire: Ω \triangleright AL AgentRequire: ω \triangleright Environment State function1: $\mathcal{L}^{(1)} \leftarrow \operatorname{seed}(\mathcal{U})$ \triangleright Create the initial labeled set2: $\mathcal{U}^{(1)} \leftarrow \mathcal{U}$ 3: for $i := 1 \dots B$ do4: $\operatorname{acc}^{(i)} \leftarrow \operatorname{Retrain}(\mathcal{L}^{(i)})$ 5: $u^{(i)} \sim \operatorname{unif}(1: |\mathcal{U}^{(i)}|)$ 6: $a^{(i)} \leftarrow \Omega(u^{(i)}) \mid a \in 1: |u^{(i)}|$ 7: $y^{(i)} \leftarrow \operatorname{label}(u_a^{(i)})$ 8: $\mathcal{L}^{(i+1)} \leftarrow \mathcal{L}^{(i)} \cup \{(u_a^{(i)}, y^{(i)})\}$ 9: $\mathcal{U}^{(i+1)} \leftarrow \mathcal{U}^{(i)} \setminus \{u_a^{(i)}\}$ 10: end for11: return $\frac{1}{B} \sum_{i=1}^{B} \operatorname{acc}^{(i)}$

Algorithm 2 Retrain			
Require: \mathcal{L}			
Require: $\mathcal{L}^{\text{test}}$			
Data			
Require: \hat{y}_{θ}			
Model			
Require: e^{\max}	▶ Maximum		
Epochs			
1: $loss^* \leftarrow \infty$)		
2: for $i := 1$.	$\dots e^{\max}$ do		
	$- heta_i - \eta abla_{ heta} \ell(\mathcal{L}, \hat{y}_{ heta})$		
	$: \operatorname{loss}_i \leftarrow \ell(\mathcal{L}^{\operatorname{test}}, \hat{y}_{\theta})$		
5: if $loss_i < loss^*$ then			
6: loss	$s^* \leftarrow loss_i$		
7: else			
8: Bre	eak		
9: end if			
10: end for			
11: return Acc	$c(\mathcal{L}^{ ext{test}}, \hat{y}_{ heta})$		

2 4 Methodology

4.1 Reproducibility

Version: 1.0

A big focus in this work is to provide an experimental setup that is fully reproducible independent from the dataset, classification model or AL algorithm used. In our opinion, an evaluation on one dataset with a given seed should always be done on the same sequence of batches x_0,\ldots,x_B . Even though different AL algorithms will pick different samples from these batches, making them unavailable for sampling in later batches, the theoretical decision tree for every possible choice in every iteration i should stay the same. This behavior is not possible with the default choice of setting a global seed at the start of the experiment, since a single additional random draw from the random number generator completely changes the decision tree for the batches. This additional random number might be drawn during the initialization of the classification model, the AL algorithm, or even during every AL iteration if Ω is stochastic. This problem also applies to the initialization of the classification model θ , the initialization and querying of Ω and the drawn mini batches during the training of θ . The desired control over all these processes can be archived by assigning a separate seeded random number generator to all these processes. To the best of our knowledge, we are the first work that discusses this issue and proposes a solution for it. We hypothesize that the insufficient

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Algorithm 3 Oracle
Require: \mathcal{U}
                                                                                                                                                       Require: \tau

    □ Unlabeled Sample Size

Require: \Omega
                                                                                                                                                                 Require: \omega
                                                                                                                                1: \mathcal{L}^{(1)} \leftarrow \operatorname{seed}(\mathcal{U})
                                                                                                                                > Create the initial labeled set
  2: \mathcal{U}^{(1)} \leftarrow \mathcal{U}
  3: for i := 1 ... B do
              acc^{(i)} \leftarrow Retrain(\mathcal{L}^{(i)})
                                                                     \triangleright \operatorname{Retrain}(\mathcal{L}^{(i)}) is shorthand for \operatorname{Retrain}(\mathcal{L}^{(i)}, \mathcal{L}^{\operatorname{test}}, \hat{y}_{\theta}, e^{\operatorname{max}})
              u^{(i)} \sim \operatorname{unif}(1:|\mathcal{U}^{(i)}|)
  5:
  6:
              r* \leftarrow -\infty
              j* \leftarrow -1
  7:
              for j := 1 \dots \tau do
  8:

    ▶ Testing every unlabeled point

                     y^{(j)} \leftarrow \text{label}(u_i^{(i)})
  9:
                     \mathcal{L}^{(j)} \leftarrow \mathcal{L}^{(i)} \cup \{(u_i^{(i)}, y^{(j)})\}
10:
                     acc^{(j)} \leftarrow Retrain(\mathcal{L}^{(j)})
11:
                     r^{(j)} \leftarrow \operatorname{acc}^{(j)} - \operatorname{acc}^{(i)}
if r^{(j)} > r^* then
12:
13:
                                                                                         ▶ Select point with largest increase in performance
                            r* \leftarrow r^{(j)}
14:
                       j* \leftarrow j  end if
15:
16:
17:
              end for
              y^{(i)} \leftarrow \text{label}(u_{i*}^{(i)})
18:
             \mathcal{L}^{(i+1)} \leftarrow \mathcal{L}^{(i)} \cup \{(u_{j*}^{(i)}, y^{(i)})\}
\mathcal{U}^{(i+1)} \leftarrow \mathcal{U}^{(i)} \setminus \{u_{j*}^{(i)}\}
19:
20:
21: end for
22: return \frac{1}{B} \sum_{i=1}^{B} acc^{(i)}
```

setup with global seeds contributes to the on-going problem of inconsistent results of AL algorithms 89 in different papers. 90

In summary, we introduce three different seeds: s_{Ω} for the acquisition function, $s_{\mathcal{D}}$ for dataset 91 splitting and mini batch sampling and s_{θ} for model initialization and sampling of dropout masks. 92 Unless stated otherwise, we will keep s_{Ω} fixed for restarts of the same experiment, while $s_{\mathcal{D}}$ and s_{θ} 93 are incremented by 1 between restarts to introduce stochasticity into our framework. 94

We compare different AL algorithms based on their median AUC score (Eq. 1) across multiple

4.2 Evaluation 95

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restarts of the experiment. This score does incorporate performance in early stages (low budget) as 98 well as capabilities to push the classifier in later stages (high budget). A good AL algorithm should be able to perform well in both scenarios. 100 Since AUC is dependent on the chosen budget, we need a general rule on how to set this hyperpa-101 rameter that does not inherently benefit a subset of algorithms. In this work, we choose the budget 102 103 per dataset to be the first point at which the oracle manages to reach a percentage of the upper bound performance. Even though we would like to propose a single percentage value for all datasets, we 104 found that different data modalities and use cases need different percentages to produce sensible 105 budgets. We propose the following values: Tabular: 99%, Image: 90% and Text: TODO. 106 Additionally, we provide evidence in Fig. 1 that previous works have not evaluated their experi-107 ments with a sufficient number of restarts. To create one slice in Fig. 1, we used all our runs for 108 the Splice dataset and drew multiple sub-sets of that respective size (X axis), effectively simulating

an experiment with that many restarts. Calculating the AUC values for every drawn run allows us

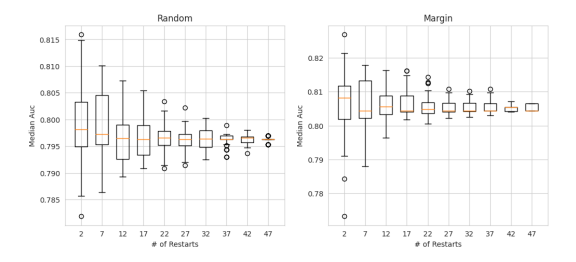


Figure 1: Random draws from Splice with different numbers of repetitions.

to draw the box-plot. Each box represents the variance of an evaluation if conducted with the respective number of restarts. We can clearly observe that low repetitions (< 10) provide an uncertain evaluation where lucky and unlucky draws of the same experiment give drastically different median AUC values. To combat is uncertain evaluation, we propose to repeat every experiment 50 times, to arrive at the true median AUC for each algorithm.

4.3 Oracle

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Version: 1.0

Posing active learning as a sequence ordering problem, the oracle sequence for a given combination 118 of dataset, model and training procedure would be the sequence that induces the highest AUC score for a given budget. However, since this combinatorial problem is not solvable for realistic datasets, 120 previous works have proposed approximations to this oracle sequence. [14] has used simulated an-121 nealing to search for the optimal sequence and used the best solution found after a fixed time budget. 122 Even though their reported performance curves display a significant lift over all other algorithms, 123 we found the computational cost of reproducing this oracle for all our datasets to be prohibitive (The 124 125 authors reported the search to take several days per dataset on 8 V100 GPUs). In this paper we propose a greedy oracle algorithm that constructs an approximation of the optimal 126 sequence in an iterative fashion. Our oracle simply tests every data point in the provided sample of 127 unlabeled points by fitting the classifier and directly measuring the resulting validation performance. 128 The point with the best validation performance is selected and added to the labeled pool for that iter-129 ation. We noticed that this oracle is overfitting on the validation set, resulting in stagnating or even 130 decreasing performance curves in later AL iterations. To circumvent this problem, we introduced 131 margin sampling as a fallback option for the oracle. Whenever the oracle does not find an unlabeled 132 point that results in an increase in performance (indicating an overfitting position), it defaults to 133 margin sampling in that iteration. The pseudocode for our oracle can be found in Alg. 3. 134

5 Implementation Details

5.1 Choosing the Classifier

Traditionally, the classifier is chosen per dataset so that it is capable of solving the dataset close to the SOTA performance reported in the literature. Similar to our hypothesis in section 4.2 we hypothesize that AL algorithms will perform similarly on small classifiers and more complex ones, so that the overall ranking of algorithms stays the same. TODO: Transform this into full hypothesis

Dataset	Architecture	Optimizer	LR	Weight Decay	Dropout
Splice	[24, 12]	NAdam	0.0012	5.9e-5	0
DNA	[24, 12]	NAdam	0.0398	3.6e-5	0
USPS	[24, 12]	Adam	0.0081	1.5e-6	0
FashionMnist	linear	-	-	-	-
Cifar10	linear	_	-	-	-

Table 2: Classifier architectures and optimized hyperparameters per dataset. Numbers in brackets signify a MLP with corresponding hidden layers.

incl. formatting?

On the basis of this hypothesis we opt to use smaller classifiers that still solve the dataset to a

reasonable degree. Smaller classifiers also require fewer labeled datapoints to acquire performance

close to the upper bound. This behavior can be observed in Figure TODO

145 For every dataset the chosen architecture's hyperparameters are optimized by to archive maximum

upper bound performance. For an overview of architectures and hyperparameters please refer to

147 Table 2.

148 5.2 Training the Classifier

149 Version: 0.1 Full rework needed

Generally, the classification model can be trained in two ways. Either you reset the parameters 150 after each AL iteration and train the classifier from scratch with the updated labeled set $\mathcal{L}^{(i)}$, or you 151 retain the previous state and only fine-tune the classifier on $\mathcal{L}^{(i)}$ for a reduced number of epochs. 152 In this work we use the fine-tuning method for raw datasets to save computation, while we use the 153 from-scratch training for embedded dataset, since they have very small classifiers and this method 154 generally produces better results. Our fine-tuning scheme always trains for at least one epoch and 155 employs an aggressive early stopping after that. The early stopping has patience 0, so it will stop as 156 soon as the validation loss does no longer decrease. 157

6 Experiments

6.1 Datasets

Version: 1.0

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For all our datasets we use the pre-defined train / test splits, if given. In the remaining cases, we

define test sets upfront and store them into separate files to keep them fixed across all experiments.

The validation set is split during experiment-time and depends on the dataset-seed.

Tabular We use **Splice**, **DNA** and **USPS** from LibSVMTools [10]. All three datasets are normalized between [0, 1].

Image We use **FashionMNIST** [13] and **Cifar10** [7]. Both datasets are normalized between [-1, 167 1].

Text We use News Category [9] and TopV2 [3]. For News Category we use the 15 most common categories as indicated by its Kaggle site. We additionally drop sentences above 80 words to reduce the necessary padding (retaining 99,86% of the data). For TopV2, we are only using the "alarm" domain. Both datasets are encoded with pre-trained GloVe embeddings [11]. Since neither set provided a fixed test set, we split random 5000 datapoints into a test set.

We would like to point out that these datasets can be considered "toy-datasets" and therefore not relevant for practical purposes. This might be true if we aimed to develop novel classification models on these datasets, however we are solely focused on comparing different AL algorithms in this paper.

Our core assumption is that a well-performing algorithm in our benchmark will transfer well into

	Splice	DNA	USPS
Oracle	0.835 ± 0.01	0.879 ± 0.01	0.870 ± 0.01
SAL_23_01_16	0.808 ± 0.01	0.857 ± 0.02	0.865 ± 0.01
Coreset_Greedy	0.808 ± 0.01	0.853 ± 0.02	0.849 ± 0.01
MarginScore	0.806 ± 0.01	0.855 ± 0.02	0.864 ± 0.01
ShannonEntropy	0.808 ± 0.01	0.855 ± 0.02	0.864 ± 0.01
RandomAgent	0.797 ± 0.01	0.823 ± 0.02	0.831 ± 0.01

	Cifar10	FashionMnist
Oracle	0.718 ± 0.01	0.721 ± 0.01
SAL_23_01_16	0.660 ± 0.01	0.667 ± 0.01
Coreset_Greedy	0.668 ± 0.01	0.674 ± 0.01
MarginScore	0.674 ± 0.01	0.677 ± 0.01
ShannonEntropy	0.671 ± 0.01	0.680 ± 0.01
RandomAgent	0.632 ± 0.02	0.633 ± 0.01

more practical use-cases.

Adapting the experimental setting from [4] we offer all our datasets in the raw setting as well as preencoded by a fixed embedding model that was trained by unsupervised contrastive learning. The text datasets are an exception, as they are only offered in their encoded form. The pre-encoded datasets offer multiple advantages in that they require smaller classification models (i.e. small MLPs) and in general much less sampled datapoints to archive close to upper bound performance (measured with the full dataset). They unify experimental setups for different datasets and therefore make them much more comparable.

185 6.2 Splice Embedded

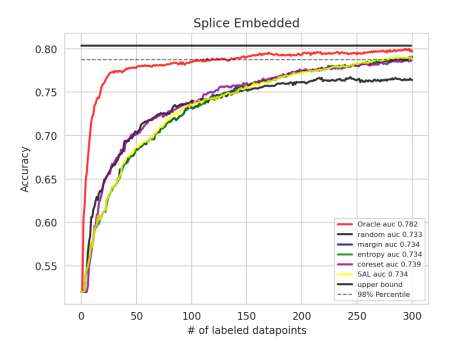


Figure 2: Results for all algorithms on the pre-encoded Splice dataset

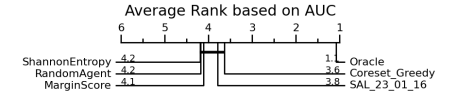


Figure 3: Critical Difference Diagram for Splice where every restart of the algorithm is one sample for the Wilcoxon-Holm method

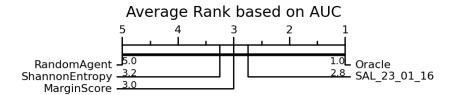


Figure 4: Critical Difference Diagram for Splice, DNA, USPS and Cifar10

7 Ablation Studies

- Setting τ to $|\mathcal{U}|$
 - Reduction of the test set for speed

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225 A Comparison of different sample sizes

