Towards Comparable Active Learning

Thorben Werner *

University of Hildesheim Universitätsplatz 1 31141 Hildesheim werner@ismll.de

Johannes Burchert*

University of Hildesheim Universitätsplatz 1, 31141 Hildesheim burchert@ismll.de

Prof. Lars Schmidt-Thieme*

University of Hildesheim
Universitätsplatz 1,31141 Hildesheim
schmidt-thieme@ismll.uni-hildesheim.de

Abstract

Active Learning has received significant attention in the field of machine learning for its potential in selecting the most informative samples for labeling, thereby reducing data annotation costs. However, the lack of reproducibility of performance gains reported in recent literature has created a chaotic landscape in Active Learning research. This paper addresses these issues of inconsistent results in active learning (AL) literature. To the best of our knowledge, we propose the first AL benchmark that tests algorithms in 3 major domains: Tabular, Image and Text. Furthermore, we highlight overlooked problems for reproducing AL experiments that can lead to unfair comparisons and increased variance in the results. To tackle these challenges, we propose an experimental protocol to accurately control the experiments. We report empirical results for 6 widely used algorithms on 7 datasets and aggregate them into a domain-specific ranking of AL algorithms.

3 1 Introduction

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Active Learning (AL) plays an important role in our society that applies machine learning to more 14 and more areas and therefore has a high demand for labeled data in more and more areas. A problem 15 that concerns academic researchers and practitioners in businesses alike and even could be extended 16 to education in schools and hobbyists around the world. On top of providing a principled way 17 to labeled unlabeled datasets, active learning is one of the two major approaches besides semi-18 supervised learning to make deep learning models more data efficient by requiring only a limited set 19 of manually labeled data. Both approaches are at their core orthogonal and can freely be combined 20 and therefore we should continue our research efforts for both approaches. 21 Among others, the authors of [18] have pointed out severe inconsistencies in results of AL papers in 22 recent years. In their supplementary materials they conducted a meta analysis of reported results of 23 several different AL algorithms and found that all considered algorithms only provided significant 24 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

^{*}Institute of Computer Science - Information Systems and Machine Learning Lab (ISMLL)

archive state-of-the-art results by significantly outperforming everyone else, while the vast majority of results proves to be non-reproducible.

30 1.1 Contributions

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- 1. Evaluation of Active Learning algorithms on datasets from 3 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.
- 3. Simple algorithm for an Oracle-Curve that can be constructed greedily and does not rely on search.

8 1.2 (From Vijaya) Problem Description

Given n=l+u data points with $l\in\mathbb{N}$ many labeled examples $\mathcal{L}=\{(x_1,y_1),\ldots,(x_l,y_l)\}$, $u\in\mathbb{N}$ many unlabeled examples $\mathcal{U}=\{x_{l+1},\ldots,x_n\}$, a budget $\mathbb{N}\ni b\leq u$ and an annotator $A:\mathbb{R}^M\to\mathbb{R}^C$ that can label x. We call $x\in\mathbb{R}^M$, $y\in\mathbb{R}^C$ predictors and labels respectively where (x,y) are drawn from an unknown distribution ρ . Find an acquisition function $F:\mathcal{U},\mathcal{L},b\mapsto\mathcal{U}_b$ that selects b many examples from \mathcal{U} such that $\mathcal{U}_b\subseteq\mathcal{U}, |\mathcal{U}_b|=b$ for labeling and the expected loss $\ell:\mathbb{R}^C\times\mathbb{R}^C\to\mathbb{R}$ (cross entropy loss) of the machine learning algorithm \hat{y} (ex. classification) trained on $\mathcal{L}\cup\{(x,A(x))\mid x\in\mathcal{U}_b\}$ is minimal:

$$\min \quad \mathbb{E}_{(x,y)\sim\rho}\ell(y,\hat{y}(x))$$

Finding the optimal subset \mathcal{U}_b is a combinatorial problem that is computationally not feasible for large b, so we allow sequential construction of the subset.

41 1.3 (From Lars) Problem Description

- Given two spaces $\mathcal{X}:=\mathcal{R}^M$ and $\mathcal{Y}:=\mathcal{R}^C$, a sample $\mathcal{D}_1,\ldots,\mathcal{D}_N\subseteq (\mathcal{X}\times\mathcal{Y})^*$ of sequences of 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

$$a: (\mathcal{X} \times \mathcal{Y})^* \times \mathcal{X}^* \to \{0,1\}^*$$

called acquisition function, s.t. the expected loss of a model learned on all predictors plus B acquired targets is minimal. Even though the acquisition function in principle could output the full subset of $\mathcal{D}_{\text{train}}$, the combinatorial problem is computationally not feasible for large B and we allow sequential construction of the subset.

2 Overview

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We constrain our work on pool-based active learning where a pool of unlabeled samples is fixed at 52 the start of each experiment and samples are chosen sequentially. Specifically, we are not experi-53 menting on so-called batch active learning, where at each iteration multiple unlabeled samples are 54 chosen at the same time. Even though batch AL is the more active research domain, it does not have 55 a principled advantage over single-sample AL except speed of computation. Not only is the problem 56 of optimizing a portfolio of unlabeled samples more complicated to solve, the algorithms also have 57 systematically less information per sample to work with. For this reason we propose to focus more 58 research effort on single-sample AL to find better algorithms in an environment that is simpler to 59 solve and easier to control. A performance comparison of batch AL and single-sample AL can be 60 found in Fig. 1, which reproduces the message of Figure 1 from the paper [5] that proposed BALD, 61 one of the SOTA algorithms for AL. We can see that the batched version of BALD [10] can at most perform on-par with the single-sample algorithm. Fig. 1 also serves as a proof of concept for our provided code base. Table 1 shows a feature comparison between our proposed benchmark and several existing benchmarks in the literature, as well as methodological AL papers with experiments on at least two data domains.

Paper	Sampling	#Datasets	#Domains	#Algorithms	Oracle
Beck et al. [2]	batch	4	1	7	-
Hu et al. [7]	batch	5	2	13	-
Li et al. [11]	batch	5	1	13	-
Zhou et al. [18]	batch	3	2	2	\checkmark
Ours	single	7	3	6	\checkmark

Table 1: Comparison of our benchmark with the existing literature. Oracle curves serve as an approximation of the best possible AL algorithm.

7 3 Related Work

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Many different algorithms have been proposed for active learning. In this work we focus on those approaches that have shown consistent results over the years as well as some of the new approaches. AL algorithms can be categorized into two classes: Geometric approaches and uncertainty-based approaches. Geometric approaches use clustering techniques to partition the data and then sample their unlabeled points based on the clusters. They often use the current classification model \hat{y}_i to encode the data into a latent space to improve the performance of their clustering. This benchmark includes the following geometric approaches: CoreSet [15], BADGE [1] and TypiClust [6]. Uncertainty-based approaches use metrics to measure the classifiers state. Commonly, a proxy for the sought after uncertainty of the model for a given datapoint is the distance of that point to the various decision boundaries, measured via the softmax output of the This benchmark includes Shannon-

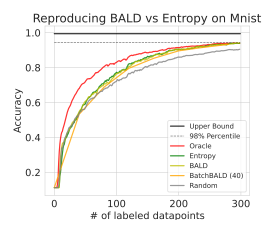


Figure 1: Shows a reproduction of the performance analysis of BALD from the original authors [5] in comparison to entropy sampling and an adaptation of BALD for batch AL [9] with a batch size of 40.

Entropy sampling [16], margin sampling [16] and BALD [9]

Some previous work also aimed to provide a benchmark suite for active learning: The authors of [2] and [11] both focus on active learning in the image domain. While [2] discuss a new metric to measure AL performance, which they call "Label Efficiency" and provide experiments on many common configurations of data preparation, model training and other hyperparameters, [11] focuses on combined approaches of AL and semi-supervised learning to aid model training. The authors of [7] study models that are learned with AL techniques in the image and text domain. They test for several different properties of the models including robustness, response to compression techniques and final performance.

99 4 Methodology

100 4.1 Evaluation

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Following [18], 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):

$$AUC(\mathcal{D}_{test}, \hat{y}, B) := \frac{1}{B} \sum_{i=1}^{B} Acc(\mathcal{D}_{test}, \hat{y}_i)$$
 (1)

where \hat{y}_i is the (re-)trained classification model after the i-th iteration. To mimic the leave-one-out protocol for cross-validation we will restart each experiment multiple times. Each restart will retain the train/test split (often given by the dataset itself), but introduces a new validation split. The AUC incorporates performance in early stages (low budget) as 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. Since AL has high variance and is prone to outliers, we propose to report median AUC instead of mean AUC values.

The AUC is dependent on the chosen budget, so we need a general rule on how to set this hyperparameter that does not inherently benefit a subset of algorithms. In this work, we choose the budget per dataset to be the first point at which any algorithm (except oracle) manages to reach a percentage of the upper bound performance measured on the full dataset. Even though we would like to propose a single percentage value for all datasets, we found that different data modalities and use cases need different percentages to produce sensible budgets. We propose the following values: **Tabular**: 99%, **Image**: 90% and **Text**: 95%.

Additionally, we provide evidence in Fig. 2 that previous works have not evaluated their experiments with a sufficient number of restarts. To create Fig. 2 we used 50 restarts from the margin/random sampling algorithm on the splice dataset. From these 50 runs we uniformly sampled subsets of runs and calculated the median AUC for this subset. One of these median AUC values corresponds to one cross-validated experiment sampled from the distribution of experiments that are restarted exactly this many times. To create one slice in Fig. 2, we drew 50 samples from this distribution. Each boxplot represents the variance of an evaluation if conducted with the respective number of restarts. We can 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 reliably arrive at the true median AUC, we propose to repeat every experiment 50 times, as only > 42 repetitions don't produce outliers anymore (as indicated by the rightmost columns in Fig 2). One way to reduce the number of necessary repetitions is to reduce the amount of variance in the experiment through specialized seeding (discussed in the next section). We ultimately decided in favor of high variance and high number of repetitions as the high variance accurately reflects real world applications of AL. Fixing for example the classifier initialization to random draws from a constant seed does not have a corresponding use-case in the real world.

4.2 Reproducibility

A big focus in this work is to provide an experimental setup that is fully reproducible independent of 135 the dataset, classification model or AL algorithm used. For a fair comparison of two AL algorithms, 136 both algorithms should receive equal starting conditions in terms of train / validation split, initialization of classifier and even the state of minor systems like the optimizer or mini-batch sampler. Even 138 though some implementations might have their own solution to some of these problems, to the best 139 of our knowledge no previous work has discussed this topic in detail. At the core of this problem 140 is the seeding utility in PyTorch, Tensorflow and other framework, whose default choice is a single global seed. Since many system draw random numbers from this seed all of them influence each 142 other to a point where a single additional draw can completely change the model initialization or data 143 split, depending on the order in the implementation. Even though you could find workarounds like re-setting the seed multiple times, this problem also extends to every AL iteration and the systems

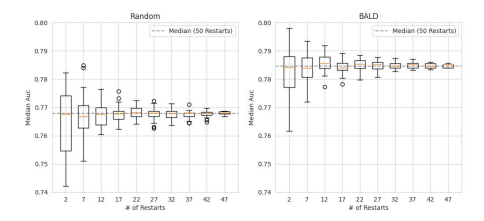


Figure 2: Random draws from an experimental distribution on the Splice dataset with different numbers of repetitions. Each point on the Y-axis represents a cross-validated result that could have been reported in a paper. This analysis shows the drastic differences in performance one could observe even when repeating an experiment 2-10 times.

within. We propose an implementation that creates a separate Random Number Generator (RNG) for each of these systems to ensure equal testing conditions even when the AL algorithm, dataset or classifier changes. We hypothesize that the insufficient setup with global seeds contributes to the on-going problem of inconsistent results of AL algorithms in different papers. In summary, we introduce three different seeds: s_{Ω} for the AL algorithm, $s_{\mathcal{D}}$ for dataset splitting and mini batch sampling and s_{θ} for model initialization and sampling of dropout masks. Unless stated otherwise, we will keep s_{Ω} fixed for restarts of the same experiment, while $s_{\mathcal{D}}$ and s_{θ} are incremented by 1 between restarts to introduce stochasticity into our framework. Some algorithms require a subsample to be drawn from \mathcal{U} in order to reduce the computational cost in each iteration, while others need access to the full unlabeled pool (i.e. for effective clustering). If a subsample is required, it will be drawn from s_{Ω} and therefore will not influence other systems in the experiments. For each algorithm, we decided if subsampling is required based on our available hardware, but decided against setting a fixed time limit per experiment, since this would introduce unnecessary

complexity into the benchmark. An overview of selected hyperparameters per AL algorithm can be

4.3 Oracle

found in Appendix C.

Posing active learning as a combinatorial problem, the oracle set \mathcal{U}_b for a given dataset, model and training procedure would be the set that induces the highest AUC score for a given budget. However, since this problem is not solvable for realistic datasets, previous works have proposed approximations to this oracle sequence. [18] has used simulated annealing to search for the optimal sequence and used the best solution found after a fixed time budget. Even though their reported performance curves display a significant lift over all other algorithms, we found the computational cost of reproducing this oracle for all our datasets to be prohibitive (The 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 sequence in an iterative fashion. Our oracle simply evaluates every data point $u_k = \text{unif}(\mathcal{U}|\mathbf{k} = 1\dots\tau)$ in a subsample of unlabeled points by fitting the classifier on $\mathcal{L}^{(i)} \cup u_k$ and directly measuring the resulting test performance. The point with the best test performance is selected and added to the labeled pool for that iteration. We noticed that this oracle is overfitting on the test set, resulting in stagnating or even decreasing performance curves in later AL iterations. This can happen for example, if the oracle picked a labeled set that enables the classifier to correctly classifier a big portion of easy samples in the test set, but now fails to find the

next single unlabeled point that would enable the classifier to succeed on one of the hard samples in the test set. This leads to a situation, where the selected data point is basically random.

To circumvent this problem, we introduced margin sampling [16] as a fallback option for the oracle. Whenever the oracle does not find an unlabeled point that results in an increase in performance, it defaults to margin sampling in that iteration. The pseudocode for our oracle can be found in Alg. 1. In the algorithm $\operatorname{Retrain}(\mathcal{L}^{(i)}, \hat{y}_{\theta})$ trains the classification model \hat{y}_{θ} and returns the accuracy on the test set $\mathcal{D}_{\text{test}}$.

 y_t is shorthand for the corresponding label of $u_t^{(i)}$ that can be recovered from the dataset labels.

When the oracle does not find a sample with positive change in classification performance $(r^* = 0)$, it applies margin sampling as a fallback $(\text{margin}(u^{(i)}, \hat{y}_{\theta}))$.

Alg. 1 replaces the algorithm in the AL process.

Algorithm 1 Oracle

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Require: \mathcal{U}, \mathcal{L}, \mathcal{Y}, \mathcal{D}_{test} Train, Margin, \tau, \hat{y}_{\theta}
   1: acc \leftarrow Train(\mathcal{L}, \mathcal{D}_{test}, \hat{y}_{\theta})
  2: r^* \leftarrow 0
  3: for t := 1 \dots \tau do
                 \mathcal{L}' \leftarrow \mathcal{L}^{(i)} \cup \{(\mathcal{U}_t, y_t)\}
                 acc' \leftarrow Train(\mathcal{L}', \mathcal{D}_{test}, \hat{y}_{\theta})
  5:
  6:
                 r \leftarrow acc - acc'
  7:
                 if r > r^* then
                         r^* \leftarrow r
  8:
  9:
                         u^* \leftarrow \mathcal{U}_t
10: if r^* = 0 then
11:
                 u^* \leftarrow \text{margin}(\mathcal{U}, \hat{y}_{\theta})
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5 Implementation Details

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At each iteration i the AL algorithm needs to pick an unlabeled datapoint based on a fixed set of information $\{\mathcal{L}^{(i)}, \mathcal{U}^{(i)}, B, |\mathcal{L}^{(i)}| - |\mathcal{L}^{(1)}|, \operatorname{acc}^{(i)}, \operatorname{acc}^{(i)}, \operatorname{opt}_{\theta}\}$, where $\theta^{(i)}$ is the current classifier and $\operatorname{opt}_{\theta}$ is the optimizer used to fit $\theta^{(i)}$. We allow algorithms to derive additional information of this set like predictions of the classifier, K-Means clustering or even training new classifiers. However, the algorithm may not incorporate external information like other datasets, queries to recover additional labels, additional training steps for the classifier, or the test/validation set.

5.1 Sampling Strategies

We selected AL algorithms that have good performances reported by multiple different sources. To ensure a fair comparison we fixed the training process of our classification model as well as the set of available information for the algorithm and selected only those that can work under these restrictions:

Uncertainty Sampling Tries to find the sample that the classifier is most uncertain about. For our benchmark we use entropy and margin (a.k.a. best-vs-second-best) sampling.

BALD [9] Applies the query-by-committee strategy of model ensembles to a single model by interpreting the model's parameters as distributions and then sample multiple outputs from them via Monte-Carlo dropout.

BADGE [1] Uses gradient embeddings of unlabeled points to select samples where the classifier is expected to change a lot. The higher the magnitude of the gradient the higher is the expected improvement in model performance.

Coreset [15] Employs K-Means clustering to try to cover the whole data distribution. Selects the unlabeled sample that is the furthest away from all cluster centers. Clustering is done in a semantically meaningful space by encoding the data with the current classifier θ_i . In this work we use the greedy variant of Coreset.

TypiClust [6] Relies on clustering similar to Coreset but proposes a new measure called "Typicality" to select unlabeled centers. Tries to select points that are in the densest regions of clusters that do not contain labeled samples yet. Clustering is done in a semantically meaningful space by encoding the data with the current classifier θ_i . It has to be pointed out that TypiClust was designed for low-budget scenarios, but we think it is still worthwhile to test and compare this algorithm with practically relevant budgets.

5.1.1 Honorable Mentions

Learning Loss for AL Introduces an updated training of the classification model with an auxiliary loss and therefore cannot be compared fairly against classification models without this boosted training regime.

214 5.2 Choosing the Classifier

Traditionally, the classifier is chosen per dataset so that it is capable of solving the dataset close to 215 the SOTA performance reported in the literature. Since we are not interested in archiving a new 216 SOTA in any classification problem, we opt to use smaller classifiers for the following reasons: 217 Smaller classifiers generally (i) exhibit more stable training behavior and (ii) on average require less 218 sampled datapoints to reach the their upper bound performance on the full dataset. For every dataset 219 the chosen architecture's hyperparameters are optimized by to archive maximum upper bound per-220 formance. One desired characteristic of small classifiers is that the ranking of AL algorithms should 221 stay the same when switching to larger models. A small analysis of this behavior can be found in 222 Appendix E. We found that the ranking of AL algorithms unfortunately does change, but we did not 223 observe systematics that benefit one or few specific algorithms. We therefore rely on the different data domains to provide classification models of different sizes and archetypes to cover all of the 225 use-cases. For an overview of architectures and hyperparameters please refer to Appendix D. 226

227 5.3 Training the Classifier

The classification model can be trained in two ways. Either you reset the parameters after each AL 228 iteration and train the classifier from scratch with the updated labeled set $\mathcal{L}^{(i)}$, or you retain the 229 previous state and only fine-tune the classifier on $\mathcal{L}^{(i)}$ for a reduced number of epochs. In this work 230 we use the fine-tuning method for raw datasets to save computation, while we use the from-scratch 231 training for embedded dataset, since they have very small classifiers and this method generally 232 produces better results. Our fine-tuning scheme always trains for at least one epoch and employs an aggressive early stopping after that. The early stopping has patience 0, so it will stop as soon as the 234 validation loss does no longer decrease. Even though the use of a fully labeled validation set might 235 be regarded as impractical, since such a set will never exist during deployment, we strongly advocate 236 for using it in benchmarks to control the classifier training. In this work we use the validation set to 237 optimize the hyperparameters of the classifier and reduce overfitting with early stopping the training 238 process in every iteration. 239

240 6 Experiments

241 6.1 Datasets

- 242 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 [13]. All three datasets are normalized between [0, 1].
- Image: We use FashionMNIST [17] and Cifar10 [10]. Both datasets are normalized according to
- their standard protocols.

 Text: We use News Category [12] and TopV2 [4]. For News Category we use the 15 most com-
- mon categories as indicated by its Kaggle site. We additionally drop sentences above 80 words to
- reduce the padding needed (retaining 99,86% of the data). For TopV2, we are only using the "alarm"
- domain. Both datasets are encoded with pre-trained GloVe (Common Crawl 840B Tokens) embed-
- domain. Both datasets are chedical with pre-trained Glove (Common Claw) 640B Tokens) embed-
- dings [14]. Since neither dataset provided a fixed test set, we randomly split 7000 datapoints into a
- test set.
- We would like to point out that these datasets can be considered "toy-datasets" and therefore not

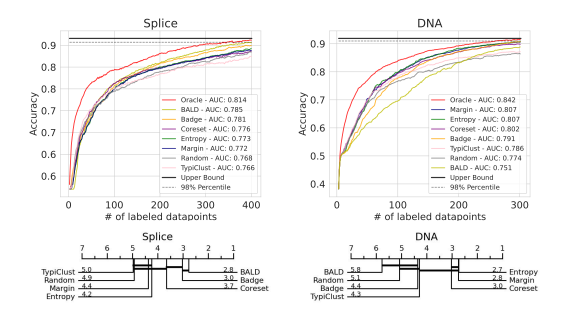


Figure 3: Results for all algorithms on Splice and DNA, both from the tabular domain. Even within one domain, the performance of the same algorithm can vary drastically.

relevant for practical purposes. This might be true if we aimed to develop novel classification models on these datasets, however, similar to our argumentation for picking smaller classifiers, 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 also transfer into more practical use-cases.

Adapting the experimental setting from [6] we offer all our datasets in the raw setting as well as pre-encoded 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 enable us to test our single-sample algorithms on more complex datasets like Cifar10 and FashionMnist without the need of sampling > 2000 datapoints before we can reach our upper bound performance. The embedding model was trained with the SimCLR [3] algorithm. For Cifar10 we adapt the reported hyperparameters from [6] and for the tabular datasets we use random search to optimize the hyperparameters. The quality of embeddings during pretext training was measured after each epoch by attaching a linear classification head and evaluating this classifier for test accuracy, mirroring our AL setup for embedded datasets.

6.2 Results

From Fig. 3 we notice drastically different qualities for the same AL algorithm for different datasets. We would like to highlight that both datasets are tabular from the medical domain with similar number of features and classes, yet we see that i.e. BALD is the best algorithm for Splice and the worst algorithm for DNA. These inconsistencies are present between the datasets of all our tested domains, further highlighting the difficulties for comparing AL algorithms in terms of average performance. In order to provide a meaningful analysis of which algorithm can be expected to perform best on average we ranked the algorithm for each dataset based on their median AUC and displayed these rankings in critical difference diagrams [8]. In Fig. 4 we report the rankings split by domain as well as across all domains (excluding the toydata).

BALD performs bad with linear classifiers since they are trained without dropout and cannot cope well with missing inputs.

TypiClust is better with embedded data not only due to lower budgets. On other datasets it is not able to outperform other algorithms in early stages

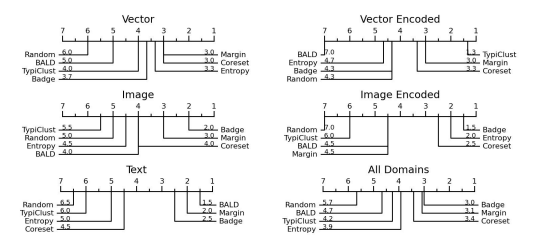


Figure 4: Critical Difference Diagram for all algorithms grouped by domain and all domains combined. Ranks are computed based on median AUC for each algorithm and dataset combination. Lower ranks are better.

284 7 Conclusion

- 285 Domains are very different
- Even within one domain we have stark differences (Fig. 3)
- 287 Best 3 ranks for all domains are 3, 3.1 and 3.4 ... No clear winner
- 288 At least on average everything is better that random

289 8 Limitations and Future Work

- 290 No batch AL
- No learned algorithms
- 292 No SOTA classifier training (data augmentation, semi-supervised, etc.)
- 293 SimCRL as Pretext task works better for images

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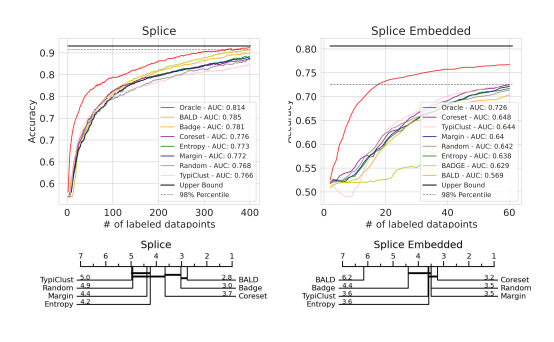
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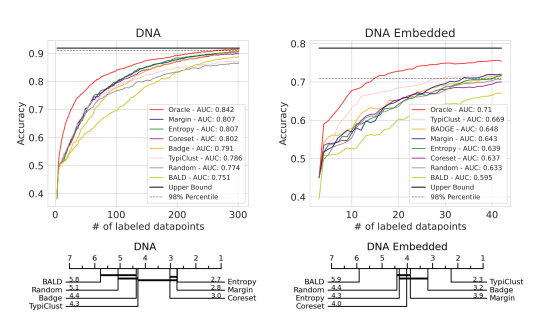
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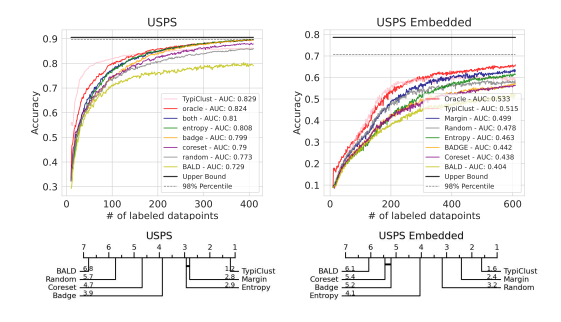
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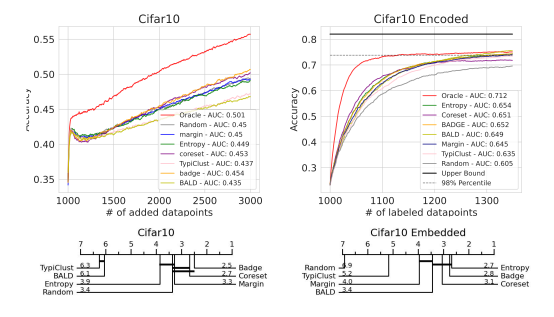
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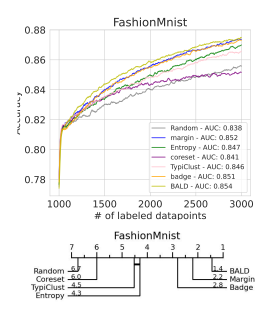
344 A All Results

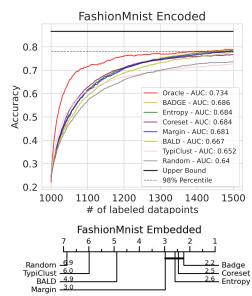


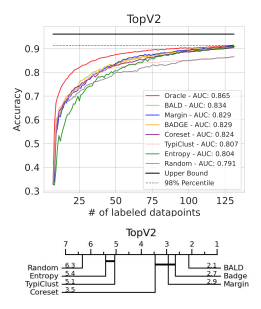


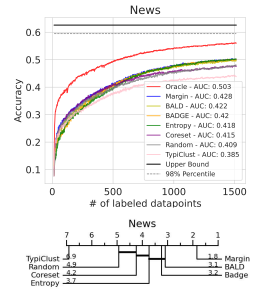




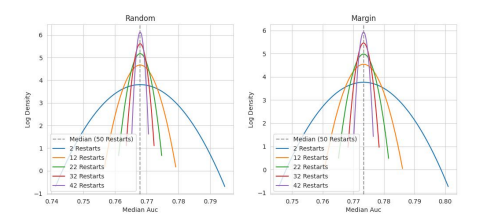








45 B Alternative Plot for Restarts Ablation



146 C Hyperparameters per AL Algorithm

TODO

350

351

347 D Hyperparameters per Dataset

Dataset	Classifier	Optimizer	LR	Weight Decay	Dropout
Splice	[24, 12]	NAdam	1.2e-3	5.9e-5	0
SpliceEnc.	linear	NAdam	6.2e-4	5.9e-6	0
DNA	[24, 12]	NAdam	3.9e-2	3.6e-5	0
DNAEnc	linear	NAdam	1.6e-3	4e-4	0
USPS	[24, 12]	Adam	8.1e-3	1.5e-6	0
USPS	linear	NAdam	7.8e-3	1.9e-6	0
FashionMnist	ResNet18	NAdam	1e-3	0	0
FashionMnistEnc	linear	Adam	1.6e-3	1e-5	5e-2
Cifar10	ResNet18	NAdam	1e-3	0	0
Cifar10Enc	linear	NAdam	1.7e-3	2.3e-5	0
TopV2	BiLSTM	NAdam	1.5e-3	1.7e-7	5e-2
News	BiLSTM	NAdam	1.5e-3	1.7e-7	5e-2

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

348 E Comparison of Different Classifier Sizes

349 We tested two different classifier sizes in Splice and DNA:

• Small: [24, 12] (2400 parameters)

• Big: [24, 48, 48] (5700 parameters)

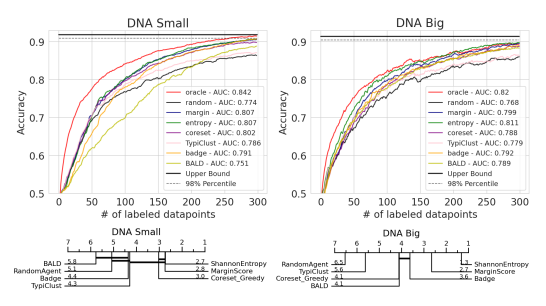


Figure 5: Comparison of small and big classifiers for the DNA dataset

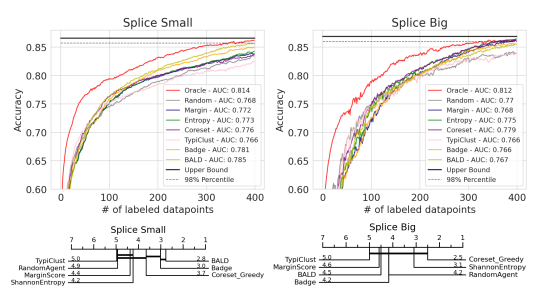
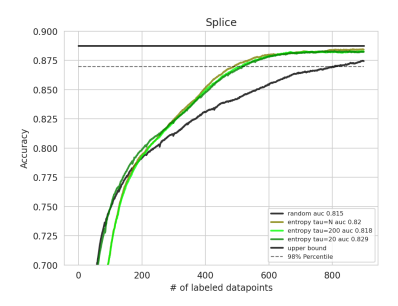


Figure 6: Comparison of small and big classifiers for the Splice dataset

F Comparison of different sample sizes



353 G AL Pseudocode

```
Algorithm 2 Active Learning
Require: U

    □ Unlabeled Pool

Require: \tau
                                                                                                                                                                   Require: \Omega
                                                                                                                                                                                                ⊳ AL Agent
Require: \omega
1: \mathcal{L}^{(1)} \leftarrow \operatorname{seed}(\mathcal{U})
                                                                                                                                                         ▶ Environment State function
                                                                                                                                                         > Create the initial labeled set
  2: \mathcal{U}^{(1)} \leftarrow \mathcal{U}
  3: for i := 1 \dots B do
4: \operatorname{acc}^{(i)} \leftarrow \operatorname{Retrain}(\mathcal{L}^{(i)})
                 a^{(i)} \leftarrow \Omega(\mathcal{U}^{(i)}) ; \quad a \in 1 : |\mathcal{U}^{(i)}|
  5:
                 y^{(i)} \leftarrow \text{label}(\mathcal{U}_a^{(i)})
  6:
  7: \mathcal{L}^{(i+1)} \leftarrow \mathcal{L}^{(i)} \cup \{(\mathcal{U}_a^{(i)}, y^{(i)})\}
8: \mathcal{U}^{(i+1)} \leftarrow \mathcal{U}^{(i)} \setminus \{\mathcal{U}_a^{(i)}\}
9: return \frac{1}{B} \sum_{i=1}^{B} \operatorname{acc}^{(i)}
```

Algorithm 3 RetrainRequire: \mathcal{L} \triangleright Labeled PoolRequire: \mathcal{D}_{val} \triangleright Validation DataRequire: \hat{y}_{θ} \triangleright Class. ModelRequire: e^{max} \triangleright Class. Model1: $loss^* \leftarrow \infty$ \triangleright Maximum Epochs2: $for i := 1 \dots e^{max} do$ \rightarrow Maximum Epochs3: $\theta_{i+1} \leftarrow \theta_i - \eta \nabla_{\theta} \ell(\mathcal{L}, \hat{y}_{\theta})$ 4: $loss_i \leftarrow \ell(\mathcal{D}^{val}, \hat{y}_{\theta})$ 5: $loss^* \leftarrow loss_i$ 7: $loss \in \mathcal{D}$ 8: $loss \in \mathcal{D}$ 9: $loss \in \mathcal{D}$ 9: $loss \in \mathcal{D}$