Boosting Active Learning via Improving Test Performance

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Abstract

Central to active learning (AL) is what data should be selected for annotation. Existing works attempt to select highly uncertain or informative data for annotation. Nevertheless, it remains unclear how selected data impacts the test performance of the task model used in AL. In this work, we explore such an impact by theoretically proving that selecting un­labeled data of higher gradient norm leads to a lower upper-bound of test loss, resulting in a better test performance. However, due to the lack of label information, directly computing gradient norm for unlabeled data is infeasible. To address this challenge, we propose two schemes, namely expected-gradnorm and entropy-gradnorm. The former computes the gradient norm by constructing an expected empirical loss while the latter constructs an unsupervised loss with entropy. Furthermore, we integrate the two schemes in a universal AL framework. We evaluate our method on classical image classification and semantic segmentation tasks. To demonstrate its competency in domain applications and its robustness to noise, we also validate our method on a cellular imaging analysis task, namely cryo-Electron Tomography subtomogram classification. Results demonstrate that our method achieves superior performance against the state of the art. We refer readers to https://arxiv.org/pdf/2112.05683.pdf for the full version of this paper which includes the appendix and source code link.

Introduction

In most scenarios, supervised learning is still the most reliable fashion for training deep neural networks. However, data annotation is often arguably expensive. To reduce the cost, active learning (AL) can be used to select a portion of all unlabeled data for annotation and the annotated data is then used to train a task model (e.g., CNN) in a supervised fashion. The goal of AL is to obtain the best test performance of the task model given a specific annotation budget. Therefore, the question becomes: what data should be selected for annotation to achieve this goal?

According to the fashion for data selection, existing AL methods can be generally categorized into two groups, namely uncertainty-based and diversity-based. The former group aims to select the most uncertain data for annotation whereas the latter group aims to select unlabeled data that diversify the labeled pool (Hasan and Roy-Chowdhury 2015; Yang et al. 2015; Mac Aodha et al. 2014; Elhamifar et al. 2013; Bilgic and Getoor 2009; Nguyen and Smeulders 2004), which can also be achieved by synthesizing new diverse labeled data (Mayer and Timofte 2020; Mahapatra et al. 2018; Zhu and Bento 2017).

Although these two main streams of methods define the state-of-the-art baselines, very little work has explored the connection between selected data and the test performance of the task model used in AL. Such a connection can be restated as: if an unlabeled sample is selected for annotation and used to train the task model, then how the test performance of the model will be impacted. Conversely, this connection can guide us to select unlabeled data which helps improve the test performance.

This work focuses on exploring the aforementioned connection. To do so, we exploit the influence function (Koh and Liang 2017) to analyze how the test performance is impacted if an individual sample is selected for annotation and used in training. Our theoretical analysis shows that selecting and annotating unlabeled data of higher gradient norm will result in a lower upper-bound of test loss, leading to a better test performance. This finding naturally leads to a data selection criterion in AL. To our best knowledge, this is the first work that theoretically analyzes how data selection impacts the test performance in AL. Moreover, to leverage our theoretical findings, we propose two independent schemes to compute the gradient norm since it cannot be directly obtained during data selection due to the lack of label information in AL settings. In the first scheme (namely expected-gradnorm), inspired by (Cai et al. 2017; Cai, Zhang, and Zhou 2013), we make use of all possible labels to compute an expected empirical loss that is adopted to compute the gradient norm. In the second scheme (namely entropy-gradnorm), we use entropy as a loss to compute the gradient norm without resorting to any label information. The rationale of the two schemes is also explained. We then propose a universal AL framework to integrate these two schemes.

Note that although we aim to boost AL via improving

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††Here, the gradient is w.r.t. model parameters rather than input.
test performance, there is no need to have an access to test data. Our theoretical analysis is completely based on the assumption that test data is unknown.

To validate our method, we conduct extensive experiments on three different tasks, including image classification, semantic segmentation, and cryo-ET (cryo-Electron Tomography) subtomogram classification (Gubins et al. 2019; Chen et al. 2017) (Appendix A.5), which is an emerging technique for analyzing the machinery of cellular systems. Our method outperforms the state-of-the-art AL baselines, demonstrating its competency and task-agnostic nature.

We summarize our main contributions as follows. Firstly, we fundamentally formulate the test performance in active learning and find that its major impact factor is gradient norm that can effectively guide unlabeled data selection. Secondly, we propose two schemes to compute gradient norm for unlabeled data without resorting to ground-truth label. Thirdly, we demonstrate that the proposed method achieves superior performance on classical computer vision challenges as well as on a domain task in computational biology.

Related Work

The Uncertainty-based AL methods rely on specialized auxiliary models to estimate uncertainty for unlabeled data. For example, (Caramalau, Bhattarai, and Kim 2021) designs a GCN (Kipf and Welling 2016) to select highly uncertain data for annotation. (Haußmann, Hannebeck, and Kandemir 2019) adopts a Bayesian network to predict uncertainty. (Zhang et al. 2020; Sinha, Ebrahimi, and Darrell 2019; Ducoffe and Precioso 2018) use adversarial learning and multiple auxiliary models (e.g. VAE (Kingma and Welling 2013)) to estimate uncertainty. (Yoo and Kweon 2019) jointly learns the task model with a loss prediction module, which aims to predict the empirical loss for unlabeled data. This loss prediction module also needs customized design and training. While the other methods do not require additional models, they still suffer from inefficient unlabeled data selection. For instance, (Sener and Savarese 2018; Kuo et al. 2018) need to solve the classical K-center or 0-1 Knapsack problem when selecting unlabeled data, and hence the time complexity is much higher than the methods (Yoo and Kweon 2019; Sinha, Ebrahimi, and Darrell 2019) which only need a feed-forward step using trained auxiliary models during data selection. Similarly, several Bayesian methods (Gal, Islam, and Ghahramani 2017; Gal and Ghahramani 2016) suffer from inefficient uncertainty estimation due to the need of thousands of feed-forward operations for each unlabeled sample.

Another category of AL methods are known as Diversity-based (a.k.a., Distribution-based). For example, (Yang et al. 2015) proposes to add a diversity regularizer in the objective function. (Elhamifar et al. 2013) uses sample diversity to provide a guidance for solving a convex optimization problem in order to select diverse data for annotation, whereas (Nguyen and Smeulders 2004) utilizes clustering to sample diverse data for annotation. Traditionally, uncertainty and diversity are treated as different criteria for data selection, however, a recent study (Loquercio, Segu, and Scaramuzza 2020) points out that they are highly correlated. In addition, with the aid of generative models (e.g. GAN (Goodfellow et al. 2014) and VAE), new data can be synthesized to diversify the labeled pool (Mayer and Timofte 2020; Mahapatra et al. 2018; Zhu and Bento 2017). Due to the difficulty and complexity in adversarial training, it needs extra efforts to deploy these methods, especially when data format is changed. For instance, the auxiliary GAN or VAE needs to be redesigned if using 3D data (e.g. cryo-ET subtomogram image, studied in Appendix A.5).

For more discussions of the related work, we refer readers to Appendix A.9, in which we discuss how our work differs essentially from (Settles, Craven, and Ray 2007; Roy and McCallum 2001).

Methodology

We firstly explore what data needs to be selected for annotation by theoretically analyzing the connection between selected data and test performance. Then we propose two schemes to compute gradient norm for unlabeled data. Lastly, we present a universal AL framework to leverage our theoretical findings and the two schemes.

What Data to be Selected

The main criterion to evaluate an AL method is the test performance of the task model. To our best knowledge, very little work has managed to interpret how selected unlabeled data directly impacts the test performance. This remains challenging since test data is normally unknown at the time of model training and data selection. In this work, we aim to draw such a connection without resorting to test data. We also aim to use this connection to guide the data selection in AL. Our ultimate goal is to select unlabeled data that makes the task model yield a better test performance.

According to (Koh and Liang 2017), we know that given a model \( f_\theta \), such as a neural network, removing a sample \( x \) from its training set will approximately influence the loss at a test sample \( x_j \) by

\[
I_{\text{loss}}(x, x_j) = \frac{1}{n} \nabla_\theta L(f_\theta(x)) \cdot H^{-1}_\theta \nabla_\theta L(f_\theta(x)),
\]

where \( n \) denotes the amount of existing training samples, and \( f_\theta(\cdot) \) refers to the feed-forward step that yields the logits output of the model \( f_\theta \), and \( H_\theta = \frac{1}{n} \sum_{i=1}^n \nabla^2_\theta L(f_\theta(x_i)) \) is the average Hessian over all training samples. For each training sample, since we want to compute its influence (if removed) on all the samples in a test dataset, we compute the total influence as follows

\[
\sum_j I_{\text{loss}}(x, x_j) = \frac{1}{n} \sum_j \nabla_\theta L(T^{c+1}(x_j)) \cdot H^{-1}_\theta \nabla_\theta L(T^{c+1}(x))
\]

where the task model in AL cycle \( c + 1 \) (i.e. \( T^{c+1} \)) is used as \( f_\theta \) in Eq. 1, and the index \( j \) is over the test set. In this section, we assume that data selection occurs in cycle \( c \), and the selected data is used in training in cycle \( c + 1 \).

As indicated in (Koh and Liang 2017), although \( I_{\text{loss}}(x, x_j) \) could be negative when a training sample is harmful to an individual \( x_j \), \( \sum_j I_{\text{loss}}(x, x_j) \) is generally positive. Intuitively, this means removing a training sample will tend to increase the expected test loss.
In AL settings, assuming the test loss of \( T^{c+1} \) is \( L^{c+1}_{\text{test}} \), if a training sample \( x \) is removed from the labeled pool and not involved in training \( T^{c+1} \), then the influenced test loss \( L^{c+1}_{\text{test}} \) can be computed by
\[
P^{c+1}_{\text{test}} = L^{c+1}_{\text{test}} + \sum_j I_{\text{loss}}(x, x_j)
\]
\[
= L^{c+1}_{\text{test}} + \frac{1}{n} \sum_j \nabla_\theta L(T^{c+1}(x_j)) \top H_\theta^{-1} H_\theta \nabla_\theta L(T^{c+1}(x)) .
\]
(3)
Since test data is unknown during data selection in AL, directly computing \( \sum_j \nabla_\theta L(T^{c+1}(x_j)) \top H_\theta^{-1} \nabla_\theta L(T^{c+1}(x)) \) is intractable. Given \( H_\theta \) is positive definite by assumption (Koh and Liang 2017), we can derive Eq. 4 by applying the Frobenius norm\(^2\) on both sides of Eq. 3, and then we have
\[
L^{c+1}_{\text{test}} = \|L^{c+1}_{\text{test}} + \frac{1}{n} \sum_j \nabla_\theta L(T^{c+1}(x_j)) \top H_\theta^{-1} \nabla_\theta L(T^{c+1}(x))\|
\leq L^{c+1}_{\text{test}} + \frac{1}{n} \|\sum_j \nabla_\theta L(T^{c+1}(x_j)) \top H_\theta^{-1} \nabla_\theta L(T^{c+1}(x))\|
\leq L^{c+1}_{\text{test}} + \frac{1}{n} \|\nabla_\theta L(T^{c+1}(x))\| \cdot \|\sum_j H_\theta^{-1} \nabla_\theta L(T^{c+1}(x_j))\|
\leq L^{c+1}_{\text{test}} + \frac{1}{n} \|\nabla_\theta L(T^{c+1}(x))\| \cdot \|\sum_j H_\theta^{-1} \nabla_\theta L(T^{c+1}(x_j))\| .
\]
(4)
For a potential test dataset, \( \|\sum_j H_\theta^{-1} \nabla_\theta L(T^{c+1}(x_j))\| \) can be regarded as a fixed term. Therefore, the upper-bound of \( L^{c+1}_{\text{test}} \) is mainly determined by \( \|\nabla_\theta L(T^{c+1}(x))\| \). However, directly computing \( \|\nabla_\theta L(T^{c+1}(x))\| \) is infeasible due to two challenges:
1. The model \( T^{c+1} \) is not available by the time of data selection in cycle \( c \);
2. \( x \) does not have a ground-truth label for computing \( L \) (addressed in next section).

Here, we address the first challenge. To do so, we involve an approximation that uses \( \|\nabla_\theta L(T^{c}(x))\| \) to bound \( \|\nabla_\theta L(T^{c+1}(x))\| \). This is reasonable since in AL, \( x \) is used to train \( T^{c+1} \) rather than \( T^c \), which means \( T^c \) has never observed \( x \) during its training. The approximation is intuitive since the gradient norm w.r.t. the model parameters of a sample (even not employed in training) is likely to decrease as the model better fits the training set. Moreover, although it is not a rigorous guarantee for any sample, we show a theoretical support that the gradient norm of the average loss is guaranteed to reduce under reasonable assumptions. The detailed proof is given in Appendix A.1. With this bound, we can derive Eq. 5 from Eq. 4.
\[
L^{c+1}_{\text{test}} \leq L^{c+1}_{\text{test}} + \frac{1}{n} \|\nabla_\theta L(T^{c+1}(x))\| \cdot \|\sum_j H_\theta^{-1} \nabla_\theta L(T^{c+1}(x_j))\|
\leq L^{c+1}_{\text{test}} + \frac{1}{n} \|\nabla_\theta L(T^c(x))\| \cdot \|\sum_j H_\theta^{-1} \nabla_\theta L(T^{c+1}(x_j))\| .
\]
(5)
\(^2\)We ignore the norm order in equations to simplify the notations.

The Eq. 5 indicates that in cycle \( c + 1 \), removing a training sample \( x \) of higher \( \|\nabla_\theta L(T^c(x))\| \) will result in a higher upper-bound of \( L^{c+1}_{\text{test}} \). Therefore, such \( x \) should be preserved in the labeled training pool of cycle \( c + 1 \). Conversely, from the perspective of cycle \( c \), during data selection, more unlabeled samples of higher \( \|\nabla_\theta L(T^c(x))\| \) should be selected for annotation and added to the labeled pool for training \( T^{c+1} \). Such a data selection scheme helps \( T^{c+1} \) maintain a lower upper-bound of \( L^{c+1}_{\text{test}} \). Therefore, we conclude that unlabeled data of higher gradient norm should be selected for annotation in AL. To better introduce our motivation of leveraging gradient norm in AL, we provide an intuitive explanation in Appendix A.3.

**Computing Gradient Norm**

The Eq. 5 can be used to guide the unlabeled data selection in AL. Specifically, we select unlabeled data \( x \) that leads to a higher \( \|\nabla_\theta L(T^c(x))\| \), in order to lower the upper-bound of the test loss, as analyzed in last section. However, computing \( \|\nabla_\theta L(T^c(x))\| \) remains challenging since computing the empirical loss \( L \) is not viable due to the lack of the label information of \( x \). To address this challenge, we propose two schemes to compute \( \|\nabla_\theta L(T^c(x))\| \), namely expected-gradnorm and entropy-gradnorm, respectively. We also analyze the rationale of the schemes in this section.

In the expected-gradnorm scheme, inspired by (Cai et al. 2017; Cai, Zhang, and Zhou 2013), we propose to use an expected empirical loss to approximate the real empirical loss. Suppose there are \( N \) classes in a given unlabeled pool, we use \( y_i \) to denote the label of the \( i^{\text{th}} \) class. Note that \( y_i \) is a label candidate of \( x \) rather than the ground-truth of \( x \). For each sample \( x \), its expected loss can be computed by
\[
L_{\text{exp}}(T^c(x)) = \sum_{i=1}^{N} P(y_i|x) L_i(T^c(x), y_i) ,
\]
(6)
where \( P(y_i|x) \) is the posterior obtained using softmax over \( T^c(x) \) and \( L_i \) is the empirical loss when the \( i^{\text{th}} \) label candidate is assumed to be the ground-truth label of \( x \). Note that this scheme does not require the real ground-truth of \( x \).

This scheme can be easily used for classification problems because the posterior \( P \) is a single vector for each individual data sample. However, for other problems, such as semantic segmentation (i.e. pixel-level classification), this scheme is not an ideal solution. It is because in this scheme we need to consider all possible labels for all individual pixels, resulting in a huge number of possibilities (see Appendix A.4 for details), which is intractable in practice. To address this challenge, we propose another scheme in next section to compute gradient norm for unlabeled data.

In the entropy-gradnorm scheme, we use the output entropy to compute gradient norm. Specifically, we use the differentiable entropy of the softmax output of the network as a loss function. Since computing entropy does not require label, this scheme is more suitable for complex tasks in which assuming label is infeasible, such as the semantic segmentation task. Following the definition of \( P(y_i|x) \) and \( N \) in Eq. 6,
the entropy loss for each sample $x$ is defined as

$$L_{ent}(T^c(x)) = -\sum_{i=1}^{N} P(y_i|x)\log P(y_i|x). \quad (7)$$

Without an access to any label information, it is reasonable to choose entropy as a surrogate of the real loss due to two reasons.

1. According to Eq. 5, adding a training sample of higher gradient norm (if computed by the entropy loss in Eq. 7) is expected to reduce the entropy of a test sample. In classification problems, reducing the entropy is usually beneficial to reducing typical supervised losses such as the cross-entropy. For example, in semi-supervised learning (Berthelot et al. 2019; Grandvalet, Bengio et al. 2005), entropy is adopted as an effective regularizer that can be minimized, naturally leading to the use of unlabeled samples in training.

2. When the task model is well trained (a good model, e.g. in later AL cycles), entropy well approximates cross-entropy (Srinivas and Fleuret 2018).

To use this scheme, one only needs to compute entropy for output posterior and back-propagate this entropy through the task model to obtain the gradient. Despite its simplicity, this scheme can be used for both classification and semantic segmentation, leading to its task-agnostic nature. Note that the model parameters will not be updated during data selection, no matter which scheme is used to compute gradient.

Note that, both the expected empirical loss defined in Eq. 6 and the entropy loss in Eq. 7 can be used in Eq. 5 as the loss function $L$, since the derivation does not require a particular form of the non-negative $L$. They also satisfy the requirements for the loss function employed in the influence function (Koh and Liang 2017) and for deriving the reduction of average gradient norm (Appendix A.1). Furthermore, we provide quantitative evaluations to validate that more than 90% of the selected samples indeed have reduced gradient norm when the loss functions defined in Eq. 6 and Eq. 7 are used to compute gradient. This further demonstrates the reliability of the approximation in Eq. 5. We show the detailed evaluations in Appendix A.2.

Proposed Active Learning Framework

To leverage our theoretical findings, we develop a universal AL framework to integrate the two schemes introduced in last section. Specifically, we train the task model with randomly selected initial annotated data and then use one of the schemes to select unlabeled data for annotation. The newly annotated data is added to the labeled pool and we re-train the task model with the updated labeled pool. Our framework makes minimal changes to the classical AL pipeline, making it easy to be implemented for different scenarios. We summarize our framework in Algorithm 1.

Experiments

To evaluate the proposed method, we conduct extensive experiments with AL settings. For fair comparison, we reproduce the baseline methods following their suggested settings and practices, such as the way of pre-processing input. In all the experiments except ImageNet, we run the AL methods for 7 cycles, corresponding to 7 different annotation budgets (i.e. from 10% to 40% with an incremental size of 5%). For the very first cycle, we randomly select 10% of the data from the unlabeled pool and use the selected data as the initial training data for all the compared methods. Then for each subsequent cycle, we select unlabeled data using a specific AL method, and re-train the task model with the updated labeled pool. Note that the term Cycle is different from Epoch, since it only corresponds to the annotation budget in AL. All the reported results are averaged over 3 runs. For ImageNet, we conduct the experiments with 5 AL cycles (i.e. annotation budget varies from 10% to 30%) and the results are averaged over 2 runs, which are acceptable for very large-scale datasets.

We compare our method with the state-of-the-art AL baselines, including random selection, core-GCN (Caramalau, Bhattarai, and Kim 2021), sral (Zhang et al. 2020), vaal (Sinha, Ebrahimi, and Darrell 2019), ll4al (Yoo and Kweon 2019), core-set (Sener and Savarese 2018), QBC (Kuo et al. 2018), and mc-dropout (Gal and Ghahramani 2016). We denote our method with exp-gn (expected-gradnorm) and ent-gn (entropy-gradnorm), respectively. We also refer readers to the Appendix for the training details (A.8), more experimental analysis (i.e. the biomedical cryo-ET experiment (A.5), the ablation study (A.6), and the time efficiency of the different methods (A.7).

Image Classification

We exploit five widely used image classification datasets in our experiments, namely Cifar10 (Krizhevsky, Hinton et al. 2009), Cifar100 (Krizhevsky, Hinton et al. 2009), SVHN
(Netzer et al. 2011), Caltech101 (Fei-Fei, Fergus, and Perona 2006), and ImageNet (Deng et al. 2009). Both Cifar datasets include 50000 training and 10000 test samples, distributed across 10 and 100 classes, respectively. SVHN also consists of 10 classes, while it includes more samples than the two Cifar datasets, namely 73257 for training and 26032 for testing. For fair comparisons with the other methods, we do not use the additional training data in SVHN. Caltech101 has 101 classes of bigger images (e.g. $300 \times 200$ pixels), which are non-uniformly distributed across the classes. Some class has up to 800 samples, while some other only includes 40 samples. ImageNet is a large-scale dataset including around 1.28 million training samples across 1000 classes. We follow the common practice to report model performance on the validation set that consists of 50000 samples.

Following the practice in (Zhang et al. 2020; Yoo and Kweon 2019), we use ResNet-18 (He et al. 2016) as the task model for all the experiments except ImageNet. Specifically, for the two Cifar and SVHN datasets, we exploit a customized version\(^3\) of ResNet18, due to the compatibility of input dimensions. For Caltech101, we exploit the original ResNet-18. To perform fair comparisons, we use the same task model for all the compared methods. For instance, (Sinha, Ebrahimi, and Darrell 2019) originally used a VGG Net as the task model, and we replace it with ResNet-18 to maintain consistency. In addition, to validate that our method is architecture-independent, we use VGG-16 (Simonyan and Zisserman 2015) as the task model for all the methods in the ImageNet task.

As shown in Fig. 1 and Fig. 2 (left), our method outperforms the baselines on all the datasets. Besides, we have the following observations that further demonstrate the advantages of our method. Firstly, for each annotation budget, our method yields a higher accuracy than the others. This is a desired property for AL methods since the annotation budget may vary in real-world scenarios. Secondly, our method needs fewer labeled samples to achieve a better performance. For example, on Cifar10, our method (ent-gn) yields an accuracy of 91.77% with 12.5K labeled samples, whereas sraal needs 2.5K more and vaal needs 5K more samples to achieve the similar performance. Thirdly, the superior performance on SVHN and Caltech101 demonstrates that our method can well handle imbalanced datasets. Fourthly, the superior performance on ImageNet demonstrates that our method is effective for very large-scale datasets. Lastly, we observe that on Cifar10 our method (ent-gn) achieves an accuracy of 94.39%, whereas training the same task model from scratch with the full dataset only yields 93.16%. This interesting finding is aligned with the observation in (Koh and Liang 2017), which suggests that some training data is harmful to neural

\[^3\]https://github.com/kuangliu/pytorch-cifar
network learning.

**Semantic Segmentation**

This experiment demonstrates the task-agnostic nature of the proposed method.

We evaluate the AL methods on the Cityscapes (Cordts et al. 2016) dataset, which is large-scale and consists of video sequences of street scenes from 50 cities. For fair comparison, we only use the standard training and validation data. Following the common practice in (Sinha, Ebrahimi, and Darrell 2019; Yu, Koltun, and Funkhouser 2017), we crop the images to a dimension of $688 \times 688$, and set the number of categories to 19.

Following the widely used settings, we adopt the dilated residual network (DRN-D-22) as the task model. We refer readers to (Yu, Koltun, and Funkhouser 2017) for the architecture details of this network.

We employ the mIoU (mean Intersection-over-Union) to measure the performance of each method. As can be seen in Fig. 2 (right), our method tends to select more unlabeled samples of higher gradient norm than the others for all the annotation budgets. In fact, semantic segmentation is a pixel-level classification task, which is more complex than image-level classification. Moreover, the Cityscapes dataset is highly imbalanced. The superior performance demonstrates that our method is robust to imbalanced datasets and suitable for complex tasks. In addition, since our method does not rely on advanced learning fashions (e.g. adversarial learning), there is no need to customize the training scheme with task change, making our method task-agnostic.

<table>
<thead>
<tr>
<th>Method</th>
<th>Budget (%)</th>
<th>10</th>
<th>15</th>
<th>20</th>
<th>25</th>
<th>30</th>
<th>35</th>
<th>40</th>
</tr>
</thead>
<tbody>
<tr>
<td>core-set (Sener and Savarese 2018)</td>
<td>217</td>
<td>230</td>
<td>284</td>
<td>339</td>
<td>329</td>
<td>289</td>
<td>255</td>
<td></td>
</tr>
<tr>
<td>vaal (Sinha, Ebrahimi, and Darrell 2019)</td>
<td>268</td>
<td>259</td>
<td>215</td>
<td>231</td>
<td>256</td>
<td>240</td>
<td>256</td>
<td></td>
</tr>
<tr>
<td>ll4al (Yoo and Kweon 2019)</td>
<td>391</td>
<td>893</td>
<td>1084</td>
<td>1710</td>
<td>1698</td>
<td>2035</td>
<td>1817</td>
<td></td>
</tr>
<tr>
<td>sral (Zhang et al. 2020)</td>
<td>506</td>
<td>819</td>
<td>1120</td>
<td>1765</td>
<td>1842</td>
<td>2133</td>
<td>2196</td>
<td></td>
</tr>
<tr>
<td>expected-gradnorm</td>
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<td>1942</td>
<td>2318</td>
<td>2427</td>
<td>2448</td>
<td></td>
</tr>
<tr>
<td>entropy-gradnorm</td>
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<td>932</td>
<td>1141</td>
<td>1993</td>
<td>2317</td>
<td>2429</td>
<td>2443</td>
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</tbody>
</table>

Table 1: Comparison of the AL methods via the number of the selected samples (out of $K = 2500$) of higher gradient norm. The Cifar10 dataset is used.

**Our Method Tends to Select More Data of Higher Gradient Norm**

Here, we investigate whether the proposed method tends to select more unlabeled samples of higher gradient norm than the AL baselines. Specifically, for unlabeled pool $\{R_U\}$, we assume that the labels $Y_U$ for all $R_U$ are temporarily available and then compute $||\nabla_{\theta}L(T(C(R_U)))||$ for each $R_U$. We sort $\{R_U\}$ by $||\nabla_{\theta}L(T(C(R_U)))||$ in ascending order to form a new set, namely $\{S_U\}$. Then we use each AL method to select a new set of unlabeled data $\{X_K\}$ from the sorted unlabeled pool $\{R_U\}$. Afterwards, we compute $\text{len}(\text{Set}), \text{Set} = \text{intersect1d}(\{S_U\}[-K:], \{X_K\}$) (in Numpy notation) and $\text{len}(\cdot)$ yields the length of a set. As shown in Table 1, for each annotation budget, our method tends to select more unlabeled samples of higher $||\nabla_{\theta}L(T(x))||$ than the others. According to the analysis in Methodology, this observation indicates that the task model trained on our selected data can yield better performance on test data, which is aligned with the goal of AL.

**Quantitative Evaluation of the Bounds in Eq. 4 and Eq. 5**

Here we quantitatively validate the bounds in Eq. 4 and Eq. 5, derived from the original target in Eq. 3, by presenting two types of evaluations.

Firstly, we evaluate how the samples selected by Eq. 5 are consistent with that selected by Eq. 3. Note that Eq. 5 is the data selection criterion used in our algorithm. Specifically, for Eq. 3 we compute the influence of each $x$ by temporar-
ly “observing” its label. At each AL cycle, we acquire the top 2500 samples selected by Eq. 3 and Eq. 5, respectively. Then we check the proportion of the overlapped samples. As shown in Table 2, more than 90% of the selected samples are consistent between the original (Eq. 3) and approximated criterion (Eq. 5). Therefore, Eq. 5 is a reasonable surrogate of Eq. 3 in AL context. Note that only in this experiment, for the purpose of evaluation, we assume test data is available in order to compute Eq. 3.

<table>
<thead>
<tr>
<th>Eq.</th>
<th>Cycle</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>2500</td>
<td>2500</td>
<td>2500</td>
<td>2500</td>
<td>2500</td>
<td>2500</td>
<td>2500</td>
<td></td>
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<tr>
<td>5</td>
<td>2290</td>
<td>2377</td>
<td>2396</td>
<td>2432</td>
<td>2449</td>
<td>2438</td>
<td>2482</td>
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</tbody>
</table>

Table 2: How the samples selected by Eq. 5 are consistent with that selected by Eq. 3, evaluated on Cifar10.

Secondly, we illustrate how the upper-bound changes after each approximation in Eq. 4 and Eq. 5. Since we actually use either Eq. 6 or Eq. 7 as the loss to compute gradient norm for unlabeled data, these upper-bounds are computed based on the proposed expected-gradnorm scheme and entropy-gradnorm scheme, respectively. Specifically, after each AL cycle, we compute four terms.

- **Target**: \( \sum \nabla_{\theta} L(T^{c+1}(x_j))^T \theta_{c+1} \nabla_{\theta} L(T^{c+1}(x_j)) \) in line 2 of Eq. 3, which is the exact influence to test loss.
- **Approx-1**: \( || \sum \nabla_{\theta} L(T^{c+1}(x_j))^T \theta_{c} \nabla_{\theta} L(T^{c+1}(x_j)) || \) in line 2 of Eq. 4.
- **Approx-2**: a further approximation in line 4 of Eq. 4: \( || \nabla_{\theta} L(T^{c+1}(x_j)) || \cdot || \sum \theta_{c} \nabla_{\theta} L(T^{c+1}(x_j)) || \).
- **Approx-3**: \( || \nabla_{\theta} L(T^{c}(x)) || \cdot || \sum \theta_{c-1} \nabla_{\theta} L(T^{c+1}(x_j)) || \) in line 2 of Eq. 5.

Note that when using the expected-gradnorm scheme, all \( L \) in above four terms will be the expected loss defined in Eq. 6. When using the entropy-gradnorm scheme, all \( L \) in the four terms will be the entropy loss defined in Eq. 7. For each term, we use all unlabeled samples to compute an average value. The results in Fig. 3 demonstrate that the bounds in the three approximations do not deviate much from the target in Eq. 3, indicating the reliability of these derived bounds.

### Better Generalization

Here, we compare the generalization of the task model that is trained on the data selected by the different AL methods. Specifically, after each AL cycle, we compute the gap between the training and test accuracy. The smaller the better for this value, since a larger one indicates that the task model suffers more from over-fitting. As shown in Table 3, our method yields the smallest gap, demonstrating its superior generalization ability.

<table>
<thead>
<tr>
<th>budget (%)</th>
<th>15</th>
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<th>25</th>
<th>30</th>
<th>35</th>
<th>40</th>
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</thead>
<tbody>
<tr>
<td>core-set</td>
<td>14.59</td>
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<td>10.88</td>
<td>9.95</td>
<td>8.92</td>
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<td>vaal</td>
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<td>12.24</td>
<td>11.08</td>
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<tr>
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<td>10.7</td>
<td>8.89</td>
<td>8.78</td>
<td>7.64</td>
<td>7.23</td>
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<tr>
<td>staal</td>
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<td>11.55</td>
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<td>8.62</td>
<td>8.01</td>
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<tr>
<td>exp-gn</td>
<td><strong>12.74</strong></td>
<td>10.04</td>
<td>8.06</td>
<td>7.05</td>
<td><strong>6.09</strong></td>
<td>5.66</td>
</tr>
<tr>
<td>ent-gn</td>
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<td><strong>9.9</strong></td>
<td><strong>7.76</strong></td>
<td><strong>6.79</strong></td>
<td>6.29</td>
<td>5.65</td>
</tr>
</tbody>
</table>

Table 3: Gap (%) between training and test accuracy after each AL cycle, evaluated on Cifar10. Annotation budget of 10% is ignored since all the methods use the same randomly selected initial data for the first cycle training.

### Conclusion

In this paper, we theoretically analyze the connection between data selection and the test performance of the task model used in active learning. We prove that selecting unlabeled data of higher gradient norm can reduce the upper-bound of the test loss. We propose two independent schemes to compute gradient norm and a universal active learning framework to leverage the schemes. We conduct extensive experiments on various benchmark datasets and the promising results validate our theoretical findings and the proposed schemes.
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References


