Out-of-Distribution Detection through Soft Clustering with Non-Negative Kernel Regression

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Abstract

As language models become more general purpose, increased attention needs to be paid to detecting out-of-distribution (OOD) instances, i.e., those not belonging to any of the distributions seen during training. Existing methods for detecting OOD data are computationally complex and storage-intensive. We propose a novel soft clustering approach for OOD detection based on non-negative kernel regression. Our approach greatly reduces computational and space complexities (up to $11\times$ improvement in inference time and 87% reduction in storage requirements). It outperforms existing approaches by up to 4 AUROC points on four benchmarks. We also introduce an entropyconstrained version of our algorithm, leading to further reductions in storage requirements (up to 97% lower than comparable approaches) while retaining competitive performance. Our soft clustering approach for OOD detection highlights its potential for detecting tail-end phenomena in extreme-scale data settings. Our source code is available on Github^{[1](#page-0-0)}.

1 Introduction

Despite the successes of generalized models of natural language, the challenge of generalization to out-of-distribution (OOD) data—data that differs from the training data distribution—remains [\(El](#page-9-0)[sahar and Gallé,](#page-9-0) [2019;](#page-9-0) [Liu et al.,](#page-9-1) [2024\)](#page-9-1). This can be a limiting obstacle in known, sensitive domains like medicine and finance [\(Yang et al.,](#page-10-0) [2023;](#page-10-0) [Salehi](#page-10-1) [et al.,](#page-10-1) [2022\)](#page-10-1), or even in "domains" which are unknown or imperceptible to humans [\(Plank,](#page-10-2) [2016\)](#page-10-2). OOD shifts are also important in detecting long tail phenomena [\(Lewis et al.,](#page-9-2) [2021;](#page-9-2) [Liu et al.,](#page-9-3) [2022\)](#page-9-3), which are critical to ensure robust and reliable application of modern language models.

Figure 1: Illustration comparing kNN (top) with kMeans (middle) and our proposed NNK-Means (bottom). We show how distance-based methods (e.g. kNN, top) may incorrectly classify OOD instances as ID if they are close to an outlier in the ID training set. Similarly, hard clustering algorithms (e.g., kMeans) may incorrectly classify ID data as OOD if it lies at the edge of a cluster. The use of soft-clustering in our method overcomes these limitations and better captures the underlying data geometry, enabling more accurate OOD detection.

While OOD detection has been extensively studied ([§2\)](#page-1-0), most approaches have limitations preventing them from being applied broadly. Existing distance-based approaches for OOD detection [\(Sun](#page-10-3) [et al.,](#page-10-3) [2022;](#page-10-3) [Breunig et al.,](#page-8-0) [2000;](#page-8-0) [Kriegel et al.,](#page-9-4) [2009\)](#page-9-4) are often not scalable as they rely on storing the entire in-distribution (ID) training set. This is particularly challenging given the size of training data for LLMs. Approaches that improve scalability make strong assumptions about the distribution of data (e.g., the ID data does not have small clusters [\(He et al.,](#page-9-5) [2003\)](#page-9-5)) or are applicable only when the data is labeled [\(Lee et al.,](#page-9-6) [2018\)](#page-9-6).

1 <https://github.com/STAC-USC/NNK-Means-OOD>

While requiring lower storage and computation, classification-based approaches for OOD detection are typically limited to cases where labeled data is available [\(Hendrycks and Gimpel,](#page-9-7) [2017\)](#page-9-7). Moreover, they perform worse than distance-based approaches [\(Liang et al.,](#page-9-8) [2018\)](#page-9-8).

In this work, we present a clustering approach for OOD detection that (i) makes no assumptions about the underlying data distribution, (ii) applies to both labeled and unlabeled data, (iii) is scalable, and (iv) is compute and storage efficient. Our OOD detection method builds on a dictionary-based approach that leverages a non-negative kernel regression (NNK)-based soft clustering technique called NNK-Means [\(Shekkizhar and Ortega,](#page-10-4) [2022\)](#page-10-4) (see [Figure 1\)](#page-0-1). Compared to hard clustering, soft clustering, i.e., associating each sample with multiple cluster centers in the data manifold, leads to a better approximation of the ID data and, consequently, improved OOD detection. It also requires fewer clusters and is, therefore, more storage-efficient. To the best of our knowledge, we are the first to leverage soft clustering for text OOD detection. We are also the first to extend the usage of NNK-Means to the text domain—introducing two novelties to do so. First, we use the approximation error as a criterion for OOD detection. Additionally, to avoid dependence on the number of cluster centers—the critical limitation in most clustering algorithms—we introduce a new, improved formulation of NNK-Means, proposing an entropy-constrained data-driven selection process. This Entropy-Constrained NNK-Means (EC-NNK-Means) also significantly reduces memory usage and inference time while maintaining comparable OOD detection performance.

We empirically validate the performance of NNK-Means for OOD detection on 4 benchmark datasets. We show that it consistently achieves superior or comparable performance relative to stateof-the-art approaches [\(Liu et al.,](#page-9-9) [2020;](#page-9-9) [Sun et al.,](#page-10-3) [2022\)](#page-10-3) while requiring over an order of magnitude lower storage and inference time. We also find that our approach is applicable across a variety of settings, effectively leveraging ID labels when they are present but providing competitive performance when they are not and maintaining high performance when using different types of embeddings. Overall, we find that our soft-clusteringbased approach yields state-of-the-art OOD detection performance while improving memory and computational efficiency–particularly when using

our improved formulation with entropy constraints.

2 Related Work

OOD detection methods in NLP broadly fall into two categories: (i) post-hoc methods that detect OOD instances after deriving their representations from pre-trained language models (PLMs) and (ii) works focused on learning representations that improve OOD detection.

Post-hoc OOD Detection These methods are typically applied to the dataset representations, which can either be *Pre-trained Representations* obtained directly from PLMs or *Fine-tuned Representations* obtained after fine-tuning the PLMs for a particular task. Post-hoc methods can be further divided into two categories. First, distancebased methods compute the minimum distance to new data from ID training data as the OOD score. For example, [Lee et al.](#page-9-6) [\(2018\)](#page-9-6) computes the class-wise Mahalanobis distance between class centroids and a query point to obtain an OOD score. [Xu et al.](#page-10-5) [\(2020\)](#page-10-5) proposed Gaussian Discriminant Analysis (GDA), which leverages Euclidean and Mahalanobis distances with generative classifiers to identify OOD instances. [Sun et al.](#page-10-3) [\(2022\)](#page-10-3) directly uses the distance to the kth nearest neighbor (KNN). However, these approaches require storing the entire ID training set, significantly increasing memory requirements. Alternatively, based on the intuition that a classifier output distribution tends to reflect training distribution, classifier-based methods leverage the output logits to get a confidence score for OOD detection. The most frequently used and simple such method uses the Maximum Softmax Probability (MSP) of the classifier as confidence, as introduced by [Hendrycks and Gimpel](#page-9-7) [\(2017\)](#page-9-7) and later improved by ODIN [\(Liang et al.,](#page-9-10) [2017\)](#page-9-10) by adding temperature scaling and input preprocessing. To tackle the over-confidence problem of MSP, [Liu et al.](#page-9-9) [\(2020\)](#page-9-9) introduces Energy, an energy-based scoring function to better detect OOD data. [Yilmaz and Toraman](#page-10-6) [\(2022\)](#page-10-6) instead proposes Distance-to-Uniform (D2U) to find the OOD data whose output distribution is closer to a uniform distribution.

Learning Representations for OOD Detection Many methods employ Supervised or Marginbased Contrastive Loss [\(Zhou et al.,](#page-11-0) [2021a\)](#page-11-0) for OOD detection, which increases the similarity of instance pairs if they belong to the same class

and decreases it otherwise. Various variants have introduced multiple improvements to enhance discrimination performance, such as Adversarial Contrastive Learning [\(Zeng et al.,](#page-10-7) [2021\)](#page-10-7), KNNenhanced Contrastive Learning (KNN-CL) [\(Zhou](#page-11-1) [et al.,](#page-11-1) [2022\)](#page-11-1), and Reassigned Contrastive Learning (RCL) [\(Wu et al.,](#page-10-8) [2022\)](#page-10-8). Apart from Contrastive Learning, [Xu et al.](#page-10-9) [\(2021\)](#page-10-9) utilizes features from all layers of PLMs to form Mahalanobis Distance Features (MDF), and GNOME [\(Chen et al.,](#page-9-11) [2023\)](#page-9-11) combines MDF from both pre-trained and fine-tuned models, while Avg-avg [\(Chen et al.,](#page-9-12) [2022\)](#page-9-12) simply averages all token representations in each intermediate layer to form the sentence representation for OOD detection.

Additionally, obtaining OOD data in real-world scenarios is challenging; thus, many methods use pseudo-OOD data for representation learning [\(Zhan et al.,](#page-10-10) [2021;](#page-10-10) [Shu et al.,](#page-10-11) [2021;](#page-10-11) [Lang et al.,](#page-9-13) [2022;](#page-9-13) [Xu et al.,](#page-10-12) [2022;](#page-10-12) [Kim et al.,](#page-9-14) [2023\)](#page-9-14). Besides these, methods like DATE [\(Manolache et al.,](#page-10-13) [2021\)](#page-10-13), PTO [\(Ouyang et al.,](#page-10-14) [2023\)](#page-10-14), and BLOOD [\(Jelenic´](#page-9-15) [et al.,](#page-9-15) [2023\)](#page-9-15) do not fit into these categories but have also achieved notable results.

Our work is a post-hoc method, which focuses primarily on techniques to detect OOD samples irrespective of the representations used. Our proposed method is computationally efficient, providing the memory benefits of clustering and classifierbased techniques while performing comparably with distance-based methods.

3 NNK-Means and Variants

We briefly present the background on soft clustering via NNK-Means [\(Shekkizhar and Ortega,](#page-10-4) [2022\)](#page-10-4) for modeling a data distribution ([§3.1\)](#page-2-0). Next, we present our extension of the method via the introduction of an entropy constraint ([§3.2\)](#page-3-0).

3.1 Background

Conventional clustering methods, such as kMeans [\(He et al.,](#page-9-5) [2003\)](#page-9-5), are trained in two steps: (i) *coding*: each training item is assigned to *one* existing cluster (corresponding to the nearest cluster center), and (ii) *dictionary update*: new cluster centers are computed, where each cluster center (dictionary atom) is the average of all training items assigned to the cluster (see [Figure 1,](#page-0-1) middle).

In contrast, a soft-clustering approach such as NNK-Means operates as follows. (i) Coding: each training item is assigned to *multiple* cluster centers (sparse coding), with non-negative weights that quantify similarity to the cluster center (larger weights for higher similarity between input and cluster center). This soft clustering allows more flexible representations with lower storage (fewer clusters can represent the data). (ii) Dictionary Update: the new cluster centers (atoms) are obtained as weighted averages of the inputs assigned to the cluster, where the weights are non-negative. The set of cluster centers is designed to minimize reconstruction error on the training data. [Figure 1](#page-0-1) (bottom) illustrates this approach.

Formally, given a dataset of N data points represented by a matrix $\mathbf{X} \in \mathbb{R}^{d \times N}$, the goal is to learn a dictionary matrix $\mathbf{D} \in \mathbb{R}^{d \times M}$ (where each column represents a cluster center) and a sparse weight matrix $\mathbf{W} \in \mathbb{R}^{M \times N}$ which generates sparse linear combinations of the columns of D that approximate the training data:

$$
\hat{\boldsymbol{D}}, \hat{\boldsymbol{W}} = \mathop{\arg\min}\limits_{\substack{\boldsymbol{D}, \boldsymbol{W}: \forall i, \boldsymbol{w}_i \geq 0, \\ \|\boldsymbol{w}_i\|_0 \leq k}} \|\boldsymbol{X} - \boldsymbol{D}\boldsymbol{W}\|_2^2 \qquad (1)
$$

Here, each column of W , w_i , is sparse, with at most k non-zero entries. To achieve this, NNK-Means alternates between *sparse coding* and *dictionary/cluster update* as follows, until a convergence criterion is reached.

Sparse Coding We find a W that minimizes reconstruction error with the current dictionary. We can rewrite the objective in [\(1\)](#page-2-1) to instead use a kernelized representation of the input data $\Phi =$ $\phi(\boldsymbol{X}) \in \mathbb{R}^{N \times N}$. Since each atom is a nonnegative linear combination of elements of Φ , the dictionary matrix can be written $\mathbf{D} = \mathbf{\Phi} \mathbf{A} \in \mathbb{R}^{d \times M}$, where $A \in \mathbb{R}^{N \times M}$ is the dictionary coefficients matrix containing the weights. Then, we can kernelize the minimization objective from [\(1\)](#page-2-1) and find each column of \hat{W} as

$$
\hat{\boldsymbol{w}}_i = \underset{\boldsymbol{w}_i \geq 0, \|\boldsymbol{w}_i\|_0 \leq k}{\arg \min} \|\boldsymbol{\phi}_i - \boldsymbol{\Phi} \boldsymbol{A} \boldsymbol{w}_i\|_2^2, \quad (2)
$$

where ϕ_i corresponds to the kernel representation of data x_i . Finding \hat{w}_i from [\(2\)](#page-2-2) involves handling an $N \times N$ kernel matrix, resulting in run times that would scale poorly with the dataset size. [Shekkizhar and Ortega'](#page-10-15)s [\(2020\)](#page-10-15) geometric insight into the NNK objective enables the efficient computation of each \hat{w}_i from a small subset of the data, specifically the k-nearest neighbors of each point. Thus, [\(2\)](#page-2-2) can be rewritten for each data point and solved with NNK as

$$
\hat{\boldsymbol{w}}_{i,S} = \underset{\boldsymbol{\theta}_i \geq 0}{\arg \min} \|\boldsymbol{\phi}_i - \boldsymbol{\Phi} \boldsymbol{A}_S \boldsymbol{\theta}_i\|_2^2 \text{ and } \hat{\boldsymbol{w}}_{i,S^c} = \mathbf{0},
$$
\n(3)

where the set S corresponds to a subset of the dictionary atoms ΦA that can have nonzero influence^{[2](#page-3-1)}. This leads to a geometric interpretation: given $\hat{w}_{i,S}$ the corresponding sparse set of selected atoms S forms a convex polytope around x_i [\(Shekkizhar](#page-10-15) [and Ortega,](#page-10-15) [2020\)](#page-10-15).

Dictionary Update Given the sparse codes W computed in the first step, this second step updates the dictionary coefficients matrix \boldsymbol{A} to minimize the reconstruction error:

$$
\mathbf{A} = \mathbf{W}^\top (\mathbf{W} \mathbf{W}^\top)^{-1}.
$$
 (4)

This update rule is similar to the Method of Optimal Directions [\(Engan et al.,](#page-9-16) [1999\)](#page-9-16) and has the advantage of keeping the cluster centers in the same space as input data.

A limitation of NNK-Means is that the number of atoms in the dictionary, M , is a hyperparameter. While dictionaries with a larger set of atoms can improve representation, they increase the complexity of coefficient selection, while also requiring more storage. In NNK-Means, there is no obvious way to adjust the number of atoms other than training the system with a new choice of M.

3.2 Entropy-Constrained NNK-Means

To address these limitations, we propose Entropy-Constrained NNK-Means (EC-NNK-Means). Our new approach estimates the number of points that select each cluster from the sparse coding weights in W . The percentage of points selecting a cluster can be viewed as a "cluster probability," which quantifies the importance of the cluster. Then, we introduce an entropy-based regularization term into the cluster optimization, which favors selecting atoms representing more data points (i.e., higher probability/lower entropy atoms).

Consider a query $q = x_i$ and the set S of its k-nearest dictionary atoms. We can expand the minimization objective in [\(3\)](#page-3-2) for each θ :

$$
\boldsymbol{\theta}_{i} = \underset{\boldsymbol{\theta} \geq 0}{\arg \min} \frac{1}{2} \boldsymbol{\theta}^{\top} \boldsymbol{K}_{S,S} \boldsymbol{\theta} - \boldsymbol{\theta}^{\top} \boldsymbol{K}_{S,q}, \quad (5)
$$

where $K_{Y,Z} = \phi(Y)^{\top} \phi(Z)$ is the chosen kernel function that encodes similarity between any given sets of vectors Y and Z.

In [\(5\)](#page-3-3), cluster assignments are influenced by the similarities between the query and its nearest cluster centers $(K_{S,q})$ and between cluster centers $(K_{S,S})$. This results in each point being assigned to a non-redundant set of its most similar atoms but does not account for the size of each cluster. The NNK-Means assignment objective can be modified to consider also the probability that a given point belongs to each cluster, represented by $p \in \mathbb{R}^M$. To do this, we include an entropy regularization term that penalizes the least selected (lower probability/higher entropy) clusters:

$$
\boldsymbol{\theta}_{i} = \underset{\boldsymbol{\theta} \geq 0}{\arg \min} \frac{1}{2} \boldsymbol{\theta}^{\top} \boldsymbol{K}_{S,S} \boldsymbol{\theta} - \boldsymbol{\theta}^{\top} \boldsymbol{K}_{S,q} + \lambda \boldsymbol{\theta}^{\top} \log p_{S},
$$
\n(6)

where p_S corresponds to the probability of each atom in the set S, and λ is a hyperparameter that controls the relative influence between the kernel similarity and probability.

The probability p_i of atom i being chosen is determined by:

$$
p_i = \frac{\sum_j \mathbb{I}(\boldsymbol{W}_{i,j} > 0)}{\sum_i \sum_j \mathbb{I}(\boldsymbol{W}_{i,j} > 0)}
$$
(7)

where $\mathbb{I}(\cdot)$ is an indicator function that is equal to 1 if the condition inside is true. This probability is defined as the number of data points assigned to atom i data with a non-zero weight normalized over the size of the dataset.

The additional entropy term added to the NNK-Means objective $(\boldsymbol{\theta}^\top \log \boldsymbol{p}_S)$ can also be regarded as the cross-entropy between the new sparse code θ and the current $\log p_S$. Minimizing this term leads to an assignment that aligns both distributions as closely as possible. Consequently, atoms that are assigned more elements during training have a higher probability of being selected by a new data point, while the reverse is true for atoms having less data assigned during training.

To adaptively learn a dictionary of a size appropriate to the data, we iteratively prune the set of M dictionary atoms to a final dictionary of size \dot{M} . Atoms with a lower probability will have fewer data points assigned in future weight assignments. When an atom's probability goes to 0, it is removed from the dictionary. This process allows for the selection of a larger initial number of atoms than the original NNK-Means, enhancing the likelihood of

²Note that we use S to denote the set to keep the notation simple, even though sets depend on the data point x_i and thus denoting them S_i would be more precise.

choosing atoms that are representative of the underlying data, while also improving efficiency by eliminating unimportant atoms. The full training procedure is described in Algorithm [1.](#page-4-0)

Algorithm 1 Entropy-Constrained NNK-Means

Input: Dataset X , training steps I , dictionary initial size N

1: $A = \{ \text{Dictionary initialized with kMeans++}^3 \}$ $A = \{ \text{Dictionary initialized with kMeans++}^3 \}$ $A = \{ \text{Dictionary initialized with kMeans++}^3 \}$ 2: $p = \left[\frac{1}{\lambda}\right]$ $\frac{1}{N}, \frac{1}{N}$ $\frac{1}{N}, \ldots, \frac{1}{N}$ $\frac{1}{N}$ $1 \times N$ 3: for iter in I do 4: $W = EC-NNK$ sparse codes 5: $p_i = \frac{\sum_j \mathbb{I}(\boldsymbol{W}_{i,j} > 0)}{\sum_i \sum_i \mathbb{I}(\boldsymbol{W}_{i,j} > 0)}$ $\overline{\sum_i \sum_j \mathbb{I}(\bm{W}_{i,j}{>}0)}$ 6: $\bm{A} = \bm{W}^\top(\bm{W}\bm{W}^\top)^{-1}$ 7: $\hat{\bm{A}} = \bm{A} \setminus \bm{A}_i, \ \forall i: p_i = 0$ 8: end for **Output:** Dictionary \hat{A} of size \hat{N}

4 NNK-Means for OOD Detection

In this section, we formally formulate the OOD detection problem ([§4.1\)](#page-4-2) and describe how to use NNK-Means for OOD detection ([§4.2\)](#page-4-3).

4.1 OOD Detection Formulation

Define an in-distribution (ID) training dataset $D_{\text{ID}} = \{(\boldsymbol{x}_1, y_1), (\boldsymbol{x}_2, y_2), \dots, (\boldsymbol{x}_N, y_N)\}\$ where x_i is a text entry and $y_i \in \{1, ..., C\}$ is the corresponding label. We also assume access to an encoder $E: \mathbf{x} \to \mathbb{R}^d$ that maps the text to a ddimensional feature space. We formulate our OOD Detection problem as a binary classification task to determine whether or not a sample is OOD with respect to the training distribution, D_{ID} , following prior work [\(Liu et al.,](#page-9-9) [2020;](#page-9-9) [Xu et al.,](#page-10-9) [2021;](#page-10-9) [Chen](#page-9-11) [et al.,](#page-9-11) [2023\)](#page-9-11). The goal is to perform a binary classification based on an OOD score $O(\mathbf{x}; E)$, where the final decision $G_{\epsilon}(\boldsymbol{x};E)$ is obtained as follows:

$$
G_{\epsilon}(\boldsymbol{x};E) = \begin{cases} \text{OOD} & \text{if } O(\boldsymbol{x};E) \leq \epsilon \\ \text{ID} & \text{if } O(\boldsymbol{x};E) > \epsilon \end{cases}, \quad (8)
$$

where ϵ represents a chosen threshold. In practice, the threshold is chosen to ensure about 95% recall.

Pipeline Our pipeline is as follows: for a given sample x , we first obtain its representations using the encoder E. These representations $E(x)$ are then passed to OOD detection methods, which can be either classifier-based or post-hoc (described in [§2\)](#page-1-0), finally yielding an OOD score, $O(·)$. The backbone model of encoder E can be a PLM or a fine-tuned version E' , which is trained on a classification task using the ID training data.

4.2 Generating OOD Scores with NNK-Means

The dictionary and assignments learned by NNK-Means are optimized to minimize the reconstruction error of the training data. New data that cannot be properly reconstructed using this dictionary, i.e., data with a higher reconstruction error, is more likely to be out-of-distribution. Therefore, we can use the definition of reconstruction error from [\(5\)](#page-3-3) as an OOD score. For any query $q \in \mathbb{R}^d$, we define its OOD score $O(q)$ as

$$
O(\boldsymbol{q}) = \frac{1}{2} \boldsymbol{\theta}_S^\top \boldsymbol{K}_{S,S} \boldsymbol{\theta}_S - \boldsymbol{\theta}_S^\top \boldsymbol{K}_{S,q} \qquad (9)
$$

Note that the value of θ is obtained by minimizing the objective in (5) , and S represents the set of k-nearest dictionary atoms to q .

We also propose C-NNK-Means, a class-wise extension incorporating label information when labeled ID data is available. Here, rather than learning one dictionary D for the entire ID dataset, we learn a separate dictionary D_c for each ID class. Then, the OOD score is:

$$
O_c(\boldsymbol{q}) = \min_c \frac{1}{2} \boldsymbol{\theta}_{S_c}^{\top} \boldsymbol{K}_{S_c,S_c} \boldsymbol{\theta}_{S_c} - \boldsymbol{\theta}_{S_c}^{\top} \boldsymbol{K}_{S_c,\boldsymbol{q}} \hspace{0.1cm} (10)
$$

For EC-NNK-Means, we set $\lambda = 0$ for the last two epochs of training and during inference. Therefore, the OOD scores for EC-NNK-Means and C-EC-NNK-Means are computed using [\(9\)](#page-4-4) and [\(10\)](#page-4-5), respectively, but using a dictionary that was learned under entropy constraints.

5 OOD Detection Experiments

5.1 Datasets

We used three datasets to empirically measure OOD detection performance: 20 Newsgroups [\(Lang,](#page-9-17) [1995\)](#page-9-17), Banking77 [\(Casanueva et al.,](#page-8-1) [2020\)](#page-8-1), and CLINC150 [\(Larson et al.,](#page-9-18) [2019\)](#page-9-18). For 20 Newsgroups and Banking77, we randomly selected 25%, 50%, and 75% of the classes to form the ID training set D_{ID} , following [Zhang et al.](#page-10-16) [\(2021\)](#page-10-16). The remaining classes were used as OOD data at test time. CLINC150 contains a designated OOD label, and the rest of the dataset was used as D_{ID} following [Lin and Gu](#page-9-19) [\(2023\)](#page-9-19). We also report results on the larger AG News [\(Zhang et al.,](#page-11-2) [2015\)](#page-11-2)

³[Arthur and Vassilvitskii](#page-8-2) [\(2007\)](#page-8-2)

in [Appendix A.](#page-11-3) Dataset statistics, splits, and other details can be found in [Appendix B.](#page-11-4)

5.2 Baselines and Models

We compared NNK-Means, our extended EC-NNK-Means, and their respective class-wise versions, C-NNK-Means and C-EC-NNK-Means, with 8 popular or recently proposed methods. For *classifier-based* OOD detection methods, we chose Maximum Softmax Probability (MSP) [\(Hendrycks and Gimpel,](#page-9-7) [2017\)](#page-9-7), Energy [\(Liu et al.,](#page-9-9) [2020\)](#page-9-9), and Distance-to-Uniform (D2U) [\(Yilmaz](#page-10-6) [and Toraman,](#page-10-6) [2022\)](#page-10-6). For *distance-based* OOD detection methods, we evaluated Mahalanobis [\(Lee et al.,](#page-9-6) [2018\)](#page-9-6) and KNN [\(Sun et al.,](#page-10-3) [2022\)](#page-10-3). We also compare against **BLOOD** (Jelenić et al., [2023\)](#page-9-15), which leverages between-layer representations, as well as kMeans and its class-wise version C-kMeans. For better illustration, we reclassified these methods into Label-Aware and Label-Blind methods, as shown in [Table 1.](#page-6-0) Label-Aware methods incorporate ID labels during training, while Label-Blind methods do not. Details of each method are provided in the [Appendix C.](#page-11-5)

We used a sentence transfomers [\(Reimers and](#page-10-17) [Gurevych,](#page-10-17) [2019\)](#page-10-17) checkpoint ^{[4](#page-5-0)} of DistilRoBERTa [\(Sanh et al.,](#page-10-18) [2020\)](#page-10-18) (82M parameters) as our encoder E. Implementation details can be found in [Ap](#page-14-0)[pendix D.](#page-14-0) [Appendix F](#page-15-0) details our hyper-parameter tuning process for some OOD detection methods.

5.3 Evaluation Metrics

We treat OOD detection as a binary classification task, where the OOD class is considered the positive sample. Following [Hendrycks and Gim](#page-9-7)[pel](#page-9-7) [\(2017\)](#page-9-7) and [Podolskiy et al.](#page-10-19) [\(2021\)](#page-10-19), we used standard evaluation metrics AUROC, AUPR, and FPR@95. We also used Inference Time (in seconds) as an additional metric to account for the efficiency of the OOD detection methods. [Appendix E](#page-15-1) provides more details.

6 OOD Detection Results and Analysis

Table [1](#page-6-0) shows the AUROC of the baselines and our proposed methods on the three evaluation datasets. AUPR and FPR@95 results are in [Appendix A.](#page-11-3)

NNK-Means outperforms baselines Overall, we find that NNK-Means and its variants have better performance than all baselines in most cases

(71% of experimental settings^{[5](#page-5-1)}). Furthermore, classifier-based approaches tend to perform worse than clustering and distance-based ones. Classifierbased approaches only had the best performance in one of the tested settings, and consistently achieved the low AUROC in all others. Despite their benefits with regards to efficiency, these approaches do not provide competitive performance.

Figure 2: Final number of atoms and AUROC for different values of Entropy Constraint hyper-parameter λ , and number of starting atoms. Reported on 20 Newsgroups with 25% ID classes. EC-NNK-Means can retain competitive performance (AUROC on top plot) with 90% less memory usage (final number of atoms on bottom plot) with $\lambda = 0.1$.

NNK-Means effectively leverages ID labels NNK-Means and kMeans are the only methods that are applicable when no labelled ID data is present, but can also incorporate label information if it is available. Nonetheless, we find that NNK-Means is better able to leverage ID labels when compared to kMeans. The label-aware variants of NNK-Means performed better than their label-blind counterparts in 71% of cases. In contrast, kMeans outperformed C-kMeans in 57% of settings. Therefore, although kMeans can incorporate ID labels, NNK-Means uses this information more effectively.

NNK-Means has low storage requirements An advantage of clustering-based methods is that the storage requirement depends on the number of clusters, not the size of the dataset. NNK-Means per-

⁴ sentence-transformers/all-distilroberta-v1

⁵In 5 out the 7 settings in [Table 1,](#page-6-0) NNK-Means and its variants have the highest AUROC.

		20 Newsgroups				Banking77		CLINIC-150
	$\%$ ID Classes \rightarrow	25%	50%	75%	25%	50%	75%	
Blind	KNN kMeans	$84.67_{\pm0.00}$ $85.17_{\pm 0.07}$	$78.83_{+0.00}$ $78.93_{\pm 0.08}$	$82.28 + 0.00$ $83.19_{+0.08}$	$94.10_{\pm 0.00}$ $94.10_{\pm0.01}$	$95.03_{\pm 0.00}$ $95.03_{\pm 0.00}$	$88.84_{+0.00}$ $88.60_{\pm0.06}$	$97.89_{\pm0.00}$ $97.88_{\pm0.02}$
abel.	$NNK-Means†$ $EC-NNK-Means^{\dagger}$	$\textbf{85.54}_{\pm 0.00}$ $85.27_{\pm 0.18}$	$79.00_{+0.07}$ $78.65_{+0.21}$	$82.23_{\pm 0.07}$ $81.86_{+0.21}$	$94.32_{\pm0.00}$ $94.27_{\pm0.01}$	$95.21_{\pm 0.02}$ $95.20_{\pm 0.01}$	$88.48_{\pm 0.03}$ $88.52_{\pm 0.06}$	$98.22_{\pm 0.02}$ $98.22_{\pm 0.03}$
Aware abel.	MSP Energy D _{2U} BLOOD Mahalanobis C-kMeans	$82.78_{\pm 0.00}$ $82.37_{+0.00}$ $83.25_{\pm0.00}$ $70.37_{+0.29}$ $80.34_{\pm 0.00}$ $85.25 + 0.06$	$74.92_{+0.00}$ $75.86_{\pm 0.00}$ $76.67_{\pm 0.00}$ $75.63_{+0.53}$ $71.33_{+0.00}$ $78.96 + 0.07$	$81.84_{\pm0.00}$ 83.94 $_{\pm 0.00}$ $83.69_{\pm0.00}$ $69.75_{\pm 0.53}$ $75.92 + 0.00$ $82.99_{+0.07}$	$88.63_{\pm0.00}$ $88.75_{+0.00}$ $88.07_{\pm0.00}$ $73.53_{\pm 0.27}$ $93.86 + 0.00$ $94.05 + 0.05$	$92.45_{\pm 0.00}$ $92.56_{\pm 0.00}$ $92.86_{\pm 0.00}$ $77.85_{\pm 0.39}$ $95.47_{+0.00}$ $95.00_{\pm 0.05}$	$85.12_{\pm 0.00}$ $83.73_{\pm0.00}$ $83.60_{\pm 0.00}$ $76.74_{\pm 0.38}$ $87.38 + 0.00$ $88.57_{\pm0.16}$	$96.43_{\pm0.00}$ $97.07_{+0.00}$ $97.15_{+0.00}$ $87.02_{\pm 0.44}$ $97.81_{\pm0.00}$ $97.89_{\pm0.00}$
	C -NNK-Means [†] C-EC-NNK-Means [†]	$85.50_{\pm 0.07}$ $85.78_{+0.15}$	$79.23_{+0.09}$ $79.20_{+0.08}$	$83.06_{+0.09}$ $83.21_{\pm 0.08}$	$94.27_{+0.05}$ $94.29_{\pm 0.08}$	$95.14_{\pm 0.04}$ $95.22_{\pm 0.04}$	$88.86_{\pm0.10}$ $88.93_{\pm0.09}$	$97.99_{+0.01}$ $98.06_{\pm0.01}$

Table 1: AUROC for OOD detection on 3 datasets with fine-tuned representations. Label-aware methods incorporate ID labels during training, while label-blind methods are unable to do so. Results are averaged over 5 random seeds. The best (↑) label-aware and label-blind methods in each column are bolded. NNK-Means and its variants, marked with †, are our methods.

		20 NG	Banking	CLINIC
	KNN	1.41	1.68	7.01
	kMeans	0.25	0.49	0.72
Label-Blind	NNK-Means [†]	0.23	0.44	0.60
	EC-NNK-Means [†]	0.23	0.40	0.59
	Mahalanobis	0.04	0.37	0.64
	C-kMeans	2.27	15.79	79.32
abel-Aware	C-NNK-Means [†]	2.27	16.68	85.67
	C -EC-NNK-Means [†]	2.24	15.51	85.89

Table 2: OOD detection Inference Time in seconds, measured on the test set and averaged over all runs for each dataset. The best (l) label-aware and labelblind methods in each column are bolded. We don't report this metric for MSP, Energy, D2U and BLOOD as explained in [Appendix E.](#page-15-1) NNK-Means and its variants, marked with †, are our methods.

forms better than all baselines while only storing 2K cluster centers instead of all 15K instances from CLINIC-150. This is 87% less storage than the best of our baselines, KNN.

[Figure 2](#page-5-2) shows how the proposed entropy constraint can reduce storage requirements even further. When working with EC-NNK-Means, the goal is to start with a large initial dictionary size and choose successively larger values of entropy-constraint hyperparameter λ until the final dictionary is of the desired size. We find that with $\lambda = 0.1$, less than 100 atoms remain in the final dictionary, but the OOD detection AUROC is comparable or better than a dictionary with 2K atoms and $\lambda = 0$. Therefore, we show that EC-NNK-Means can achieve comparable or better performance than NNK-Means and KNN while using 95% and 97% less memory, respectively. This reduced memory requirement is particularly useful when working with large datasets - where storing and running computations on the entire ID train set may be challenging.

NNK-Means has reduced inference time [Ta](#page-6-1)[ble 2](#page-6-1) shows that NNK-Means is significantly faster than KNN as operating on the smaller, learned dictionaries is quicker than working with the entire ID train dataset. In particular, on the CLINIC-150 dataset, EC-NNK-Means provides an $11\times$ reduction in inference time relative to KNN. Class-wise variants of NNK-Means have higher inference time because they involve iterating through one dictionary per ID class, an operation that is not parallelized like the computations in NNK-Means.

Figure 3: 2D visualization of 20 Newsgroups validation dataset and learned clusters, with 25% ID classes.

Table 3: Example of OOD instances overlapping with ID data from the visualization in [Figure 3,](#page-6-2) with identical label colors. Last column represents the NNK-Means Error, as presented in [\(9\)](#page-4-4). All ID and OOD instances mention the purchase or sale of a product, despite belonging to different classes. Bolded text is edited from the original to preserve anonymity.

Figure 4: OOD Detection AUROC on 20 Newsgroups with 50% ID classes, using 3 different types of representations. Results are averaged over 5 random seeds. In the first legend, different hues represent different types of methods, including baselines and our methods in both Label-Blind and Label-Aware settings. For the second legend, two shades of the same hue distinguish between methods of the same type. For example, C-EC-NNK-Means represented by dark purple and C-NNK-Means represented by light purple are both Label-Aware methods we proposed.

Competitive performance with different embeddings A key benefit of NNK-Means is its applicability in various settings, independent of the embeddings being used. To empirically validate the performance of our methods when using different representations, we evaluate OOD detection performance using two different types of embeddings, as presented in [Table 8.](#page-13-0) We report results on the 20 Newsgroups dataset, comparing pre-trained embeddings and embeddings from the same encoder model but fine-tuned with margin-based contrastive loss, as in [Zhou et al.](#page-11-6) [\(2021b\)](#page-11-6).

We find that NNK-Means provides competitive performance, outperforming all baselines even when different representations are used (see [Fig](#page-7-0)[ure 4\)](#page-7-0). In particular, when using pre-trained representations, NNK-Means performs significantly better than all other baselines (4 AUROC points better than best baseline, KNN). [Appendix A](#page-11-3) provides further results with different types of embeddings.

Qualitative analysis of clustering [Figure 3](#page-6-2) uses UMAP [\(McInnes et al.,](#page-10-20) [2020\)](#page-10-20) to visualize the results of our clustering process. We find that our clustering works as expected: when a dictionary learned on the training set is used to cluster the validation data, instances with the same class label are assigned to the same clusters. We also see separate clusters of OOD data when their class labels are substantially different from the ID labels. In some cases, there is overlap between OOD instances and ID data, such as the blue "misc.forsale" class. Analysing the text of these OOD documents shows that this overlap occurs because the OOD

and ID instances are similar (see [Table 3\)](#page-7-1).

NNK-Means' OOD score is correlated with downstream performance In [Figure 5](#page-8-3) we present the distribution of NNK-Means OOD scores for correctly and incorrectly classified instances in the Banking77 test dataset. We find that incorrectly classified instances tend to have higher OOD scores, with a median score of 0.149 for misclassified instances and 0.025 for correctly classified ones. This result highlights the advantage of OOD detection because higher OOD scores are indicative of worse downstream task performance. So, by computing this score, it is possible to identify instances where the model is likely to perform poorly and use this information to refine the training set.

Figure 5: NNK-Means OOD score for misclassified and correctly classified instances. Reported using finetuned representations on the Banking77 dataset with 75% known classes. Misclassified instances tend to have a higher OOD score than correctly classified ones.

7 Conclusion

We address the problem of OOD detection using NNK-Means, a soft-clustering algorithm. NNK-Means achieves state-of-the-art performance across 4 benchmark datasets, while requiring lower storage and improving computational efficiency relative to previous approaches that perform comparably. We introduce EC-NNK-Means, an extension of NNK-Means, and show that it can lead to further improvements in efficiency while matching or improving OOD detection performance. Our methods provide competitive performance regardless of the availability of labels or the type of embeddings used, and yield intuitive clustering of input data. Future work will explore applying our algorithms

to analyze large pre-training datasets.

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Ethical Considerations

Our work aims to enable the robust and reliable deployment of Language Models by appropriately flagging OOD data and preventing inaccurate or unpredictable output. We do not anticipate any risks or harmful consequences stemming from our work. Our code and models are publicly available to ensure our work is reproducible. All datasets used in this paper are publicly available.

Limitations

There is a multitude of approaches for OOD detection, however, we were only able to compare against a subset of these approaches. Furthermore, our datasets, models, and experiments are all English-only. Finally, our experiments used data from classes that were unseen during training to simulate OOD data. In practice, there are many different ways a system may encounter OOD instances, and our experiments may not have covered them all.

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A Additional Results

We provide additional results on AUPR (see [Ta](#page-12-0)[ble 4\)](#page-12-0) and FPR@95 (see [Table 5\)](#page-12-1) for the 3 main datasets aligned with [Table 1.](#page-6-0) To demonstrate that our methods perform relatively better on larger datasets, we also include results on AG News; see [Table 7](#page-13-1) for more details. Additionally, to show the competitive performance of our proposed methods with different representations (detailed analysis in Section [6\)](#page-5-3), we also provide the AUROC results with Pre-Trained Embeddings and Marginbased Contrastive Loss Embeddings (see [Table 8\)](#page-13-0), which are reported for 50% ID classes ratio using label-blind and label-aware methods on 20 Newsgroups.

B Datasets

In this section, we specifically introduce the four datasets we used and how they were partitioned. Each dataset was divided into training/validation/test sets. In [Table 6,](#page-12-2) we provide the statistical details of these datasets before distinguishing between ID and OOD classes.

20 Newsgroups [\(Lang,](#page-9-17) [1995\)](#page-9-17) 20 Newsgroups is a widely used benchmark for text classification, consisting of approximately 18000 newsgroup documents organized into 20 classes, each representing a specific topic such as politics, religion, science, and technology. We utilized the 20 Newsgroups dataset provided by scikit-learn and removed headers, signature blocks, and quotation blocks respectively as suggested. Following [Zhou et al.](#page-11-6) [\(2021b\)](#page-11-6), we divided the whole dataset into training/validation/test sets in an 80/10/10 ratio using stratified sampling based on class labels. For the training set, we randomly selected 25%, 50%, and 75% of the classes as ID classes and removed the remaining classes, resulting in the dataset D_{IN} . In the validation and test sets, these selected classes were considered as IN class during the OOD detection phase, while the other classes were treated as OOD class.

Banking77 [\(Casanueva et al.,](#page-8-1) [2020\)](#page-8-1) Banking77 is a specialized dataset for intent classification in the banking domain. It consists of 13083 customer service queries categorized into 77 distinct classes, each representing a specific banking-related intent. We used the HuggingFace version of this dataset, which includes 10003 user queries in the training set and 3080 queries in the test set. We split its training set into training and validation sets in a 90/10 ratio and applied the same preprocessing steps to the training set as we did with the 20 Newsgroups.

CLINC150 [\(Larson et al.,](#page-9-18) [2019\)](#page-9-18) CLINC150 is a dataset tailored for OOD intent detection. It includes 150 distinct intent classes from various domains and one designated OOD class for evaluation. The dataset consists of a total of 22500 ID queries and 1200 OOD queries. We used the ID training data directly as our training set and combined the ID validation and test data with the OOD validation and test data to form our validation and test sets, respectively.

AG News [\(Zhang et al.,](#page-11-2) [2015\)](#page-11-2) AG News is a topic classification dataset collected from various news sources, encompassing a total of four topics. We used the HuggingFace version of this dataset, which includes 120000 entries in the training set and 7600 entries in the test set. We extracted 6% of the training data to form a validation set. When selecting 25% of the classes as ID classes, AG News only includes one class, making it unsuitable for classification tasks. Therefore, we only used the 50% and 75% settings for our experiments. The rest of the processing is similar to that applied to the 20 Newsgroups dataset.

C Baselines and Models

In this section, we provide a more detailed introduction to our baselines. Mathematical notations follow the conventions established in Section [4.1.](#page-4-2)

		20 Newsgroups				Banking77			
	$\%$ ID Classes \rightarrow	25%	50%	75%	25%	50%	75%	CLINIC-150	
-Blind	KNN	$55.48_{+0.00}$	$75.56_{\pm0.00}$	$87.96_{\pm0.00}$	$87.15_{\pm 0.00}$	$95.22_{\pm 0.00}$	$95.65_{\pm 0.00}$	$99.47_{\pm0.00}$	
	kMeans	$56.00_{+0.30}$	$75.04_{+0.19}$	$87.90_{\pm 0.08}$	$87.12_{\pm 0.05}$	$95.24_{\pm 0.05}$	$95.51_{\pm 0.06}$	99.46 $_{\pm 0.01}$	
Label-	$NNK-Means†$	$56.66 + 0.00$	$75.34_{\pm 0.08}$	$87.90_{\pm 0.07}$	$87.38_{\pm0.00}$	$95.36_{\pm0.02}$	$95.55_{\pm 0.02}$	$99.54_{\pm0.01}$	
	$EC-NNK-Means$ ^T	$56.38_{\pm0.25}$	$74.95 + 0.22$	$87.78 + 0.06$	$87.28 + 0.03$	$95.31_{\pm 0.02}$	$95.57_{\pm0.04}$	$99.54_{\pm0.01}$	
Aware abel.	MSP Energy D ₂ U BLOOD Mahalanobis C-kMeans	$66.23_{\pm0.00}$ $59.84_{+0.00}$ $66.53_{+0.00}$ $42.42_{\pm 0.34}$ $51.51_{+0.00}$ $55.87_{+0.20}$	$77.71_{+0.00}$ $76.87_{+0.00}$ $78.32_{\pm0.00}$ $78.72_{+0.64}$ $69.89_{+0.00}$ $75.61_{\pm0.10}$	$89.11_{+0.00}$ $90.84_{\pm0.00}$ $90.73_{\pm 0.00}$ $88.78_{\pm 0.25}$ $84.73_{\pm 0.00}$ $87.91_{\pm 0.04}$	$70.56_{+0.00}$ $71.79_{\pm 0.00}$ $68.80_{\pm 0.00}$ $43.50_{+0.93}$ $85.82_{+0.00}$ $86.64_{\pm0.30}$	$92.90_{\pm 0.00}$ $92.87_{\pm0.00}$ $93.37_{\pm0.00}$ $77.18_{+0.52}$ $95.38_{\pm0.00}$ $95.18_{\pm 0.07}$	$91.93_{\pm 0.00}$ $92.03 + 0.00$ $91.92_{\pm0.00}$ $89.28_{\pm 0.26}$ $94.94 + 0.00$ $95.46_{\pm0.10}$	$99.01_{\pm 0.00}$ $99.19_{\pm0.00}$ $99.21_{+0.00}$ $96.37_{\pm0.14}$ $99.44_{\pm0.00}$ $99.47_{\pm0.00}$	
	C -NNK-Means [†]	$56.24_{\pm 0.23}$	$75.51_{\pm 0.10}$	$87.88_{\pm 0.06}$	$86.82_{\pm0.21}$	$95.28_{\pm 0.05}$	$95.66_{\pm0.06}$	$99.50_{\pm0.01}$	
	C -EC-NNK-Means [†]	$56.80_{+0.61}$	$75.45_{\pm 0.19}$	$87.85_{\pm0.11}$	$86.43_{\pm 0.30}$	$95.33_{\pm0.02}$	$95.68_{\pm0.07}$	$99.51_{\pm0.01}$	

Table 4: AUPR for OOD detection on 3 datasets with fine-tuned representations. Label-aware methods incorporate ID labels during training, while label-blind methods are unable to do so. Results are averaged over 5 random seeds. The best (↑) label-aware and label-blind methods in each column are bolded. NNK-Means and its variants, marked with †, are our methods.

		20 Newsgroups				Banking77				
	$\%$ ID Classes \rightarrow	25%	50%	75%	25%	50%	75%	CLINIC-150		
abel-Blind	KNN kMeans	$50.92_{\pm 0.00}$ $48.69_{+1.17}$	$70.61 + 0.00$ $71.41_{\pm 0.30}$	$82.53 + 0.00$ $81.73_{\pm 0.87}$	$32.07 + 0.00$ $32.01_{\pm 0.02}$	$25.26 + 0.00$ $25.32_{+0.01}$	$45.88_{\pm0.00}$ $46.63_{\pm 0.04}$	$10.90_{\pm 0.00}$ $10.72_{\pm 0.26}$		
	$NNK-Means†$ $EC-NNK-Means$ ^T	$50.64_{\pm 0.00}$ $52.24_{\pm 1.97}$	$71.43_{\pm 0.49}$ $72.36 + 0.44$	$78.99_{+0.79}$ $79.07_{\pm 0.51}$	$27.67_{\pm0.00}$ $27.84_{\pm0.02}$	$25.30_{+0.03}$ $25.53_{\pm 0.02}$	$50.05 + 0.01$ $49.93_{\pm0.02}$	$8.48_{\pm 0.41}$ $8.42_{+0.29}$		
Aware abel.	MSP Energy D _{2U} BLOOD Mahalanobis C-kMeans	$77.23_{\pm 0.00}$ $77.37_{+0.00}$ $76.10_{+0.00}$ $87.12_{+0.68}$ $68.32_{\pm 0.00}$ $49.12_{+1.78}$	$87.31_{\pm 0.00}$ $87.42_{\pm 0.00}$ $87.64_{\pm0.00}$ $92.10_{+0.74}$ $86.98_{\pm0.00}$ $71.54_{+0.71}$	$90.31_{\pm 0.00}$ $87.79_{\pm0.00}$ $87.16_{\pm 0.00}$ $90.36_{\pm 0.49}$ $92.21_{\pm 0.00}$ $81.90_{\pm 0.46}$	$39.35 + 0.00$ $40.04_{\pm 0.00}$ $39.27_{\pm 0.00}$ $70.82_{\pm 1.72}$ $30.22_{\pm 0.00}$ $30.10_{+0.63}$	$48.01_{\pm 0.00}$ $39.68 + 0.00$ $39.94_{\pm 0.00}$ $71.89_{+1.17}$ $23.46_{\pm 0.00}$ $24.92_{+0.25}$	$58.13_{\pm 0.00}$ $61.25_{\pm 0.00}$ $61.13_{+0.00}$ $72.58_{+1.45}$ $52.50_{+0.00}$ $47.03_{\pm 0.34}$	$17.00_{\pm 0.00}$ $12.90_{\pm 0.00}$ $12.70_{+0.00}$ $58.12_{+1.01}$ $9.60_{+0.00}$ $10.90_{\pm 0.00}$		
	C -NNK-Means [†] C -EC-NNK-Means [†]	$50.11_{\pm 0.66}$ $47.43_{\pm 0.91}$	$71.09_{+0.74}$ $71.74_{\pm 0.49}$	$80.93_{+0.25}$ $81.31_{\pm 0.28}$	$31.03_{\pm 0.47}$ $29.96_{\pm 1.48}$	$24.40_{\pm 0.40}$ $24.14_{\pm 0.44}$	$46.68 + 0.33$ $46.90 + 0.56$	$10.22_{\pm 0.13}$ $9.92_{\pm 0.28}$		

Table 5: FPR@95 for OOD detection on 3 datasets with fine-tuned representations. Label-aware methods incorporate ID labels during training, while label-blind methods are unable to do so. Results are averaged over 5 random seeds. The best (\downarrow) label-aware and label-blind methods in each column are **bolded**. NNK-Means and its variants, marked with †, are our methods.

Dataset	# Training	# Validation	# Test	# Classes
20 Newsgroups	15076	1885	1885	20
Banking77	9002	1001	3080	77
CLINC150	15000	3100	5500	$150+1$
AG News	112800	7200	7600	

Table 6: Dataset summary with statistical details about the training, validation, and test sets along with the number of classes. Note that the number of training examples is initial.

Maximum Softmax Probability (MSP) [Hendrycks and Gimpel](#page-9-7) [\(2017\)](#page-9-7) propose this method as the baseline for detecting OOD examples which has been widely adopted. For MSP, $O(\mathbf{x}; E')$ is the maximum softmax probability among any of the classes:

$$
O(\boldsymbol{x}; E') = \max_{c \in \{1, \dots, C\}} p_c(E'(\boldsymbol{x})) \qquad (11)
$$

where $p_c(\cdot)$ refers to the softmax probability for class c. Note that this method is applicable only when using fine-tuned encoder E' .

Energy [Liu et al.](#page-9-9) [\(2020\)](#page-9-9) introduces the free energy function to detect OOD samples, which can replace the Softmax Confidence Score to avoid the overconfidence problem of the softmax function. The ID data tends to have low energy scores while OOD data tends to have high scores. The free energy function is formulated as follows:

$$
\text{Energy}(\boldsymbol{x}) = \sum_{i=1}^{C} e^{f_i(E'(\boldsymbol{x}))} \tag{12}
$$

where $f_i(\cdot)$ represents the output logits for the *i*th class, and C is the number of all classes. The score $O(\mathbf{x}; E')$ is then defined as the negative of

		AUROC (\uparrow)		AUPR (\uparrow)		$\text{FPR} @ 95 (\downarrow)$		Infer. Time (\downarrow)	
	$\%$ ID Classes \rightarrow	50%	75%	50%	75%	50%	75%		
	KNN	83.75	93.09	83.03	97.23	46.47	31.71	18.50	
	kMeans	83.54	93.49	82.74	97.30	47.35	27.61	0.89	
abel-Blind	$NNK-MeansT$	83.91	93.22	82.70	97.30	45.33	30.45	1.44	
	$EC-NNK-Means^{\dagger}$	84.07	93.43	83.64	97.31	46.01	28.36	0.95	
Label-Aware	MSP Energy D _{2U} BLOOD Mahalanobis C-kMeans	82.84 79.90 82.84 77.95 83.42 83.72	86.07 86.68 87.72 86.16 92.10 93.42	83.97 79.01 84.06 75.35 83.79 82.96	94.78 94.81 95.25 93.73 96.79 97.25	53.58 55.13 53.56 53.62 53.54 47.59	55.80 46.68 46.68 51.45 34.08 27.74	0.02 2.03	
	C -NNK-Means [†]	83.26	93.37	82.20	97.31	46.92	28.72	2.18	
	C -EC-NNK-Means ¹	86.30	94.47	86.47	97.98	45.87	26.80	1.98	

Table 7: OOD detection performance on AG News are reported for AUROC, AUPR, FPR@95 and Inference Time in seconds with fine-tuned representations. Label-aware methods incorporate ID labels during training, while label-blind methods are unable to do so. Results are averaged over 5 random seeds. The best label-aware and label-blind methods in each column are bolded. We do not report Inferece Time for MSP, Energy, D2U, and BLOOD as discussed in [Appendix E.](#page-15-1) NNK-Means and its variants, marked with †, are our methods.

		Pre-Trained			Margin-based Contrastive Loss			
	$\%$ ID Classes \rightarrow	25%	50%	75%	25%	50%	75%	
	KNN	71.75	69.11	67.63	79.95	77.50	78.82	
	kMeans	70.35	68.59	67.43	80.32	78.21	80.07	
Label-Blind	$NNK-Means^{\dagger}$	75.52	72.49	69.52	80.27	77.70	79.10	
	$EC-NNK-Means^{\dagger}$	75.01	70.42	67.82	80.15	77.60	79.01	
Label-Aware	MSP Energy D2U BLOOD Mahalanobis C-kMeans	62.75 70.29	58.77 68.56	$\overline{}$ $\qquad \qquad \blacksquare$ $\qquad \qquad -$ $\qquad \qquad \blacksquare$ 57.64 68.45	78.26 78.55 79.19 69.02 76.07 80.29	75.59 75.46 76.37 73.64 74.08 78.27	77.36 78.42 78.77 65.74 72.91 79.73	
	C -NNK-Means [†]	75.08	72.76	70.48	80.42	78.12	79.67	
	C -EC-NNK-Means ¹	76.49	71.71	70.20	80.53	78.42	79.55	

Table 8: AUROC comparison with Pre-Trained Embeddings and Margin-based Contrastive Loss Embeddings. Results are reported for 50% ID classes ratio using label-blind and label-aware methods on 20 Newsgroups. The best (†) label-aware and label-blind methods in each column are **bolded**. We do not report Pre-Trained Embedding results for MSP, Energy, D2U, and BLOOD as discussed in [Appendix C.](#page-11-5) NNK-Means and its variants, marked with †, are our methods.

the energy:

$$
O(\boldsymbol{x}; E') = -\text{Energy}(\boldsymbol{x}) \tag{13}
$$

Note that this method is also applicable only when using fine-tuned encoder E' .

Distance to Uniform (D2U) Based on the idea that output distributions of OOD samples get closer to the uniform distribution than that of ID samples, [Yilmaz and Toraman](#page-10-6) [\(2022\)](#page-10-6) introduces Distanceto-Uniform (D2U), which utilizes the shape of the entire output distribution and calculates its distance to the uniform distribution as a metric to evaluate the likelihood of an example being OOD:

$$
O(\boldsymbol{x}; E') = \text{dst}(\boldsymbol{p}(E'(\boldsymbol{x})), U) \tag{14}
$$

where $p(\cdot)$ is the output softmax distribution and U refers to the uniform distribution. We follow [Yilmaz and Toraman](#page-10-6) [\(2022\)](#page-10-6)'s setting to use the KL divergence as the distance function. Note that this method is also applicable only when using finetuned encoder E' .

BLOOD The BLOOD score proposed by Jelenić [et al.](#page-9-15) [\(2023\)](#page-9-15) is a method for detecting OOD data in Transformer-based models by examining the smoothness of transformations between intermediate layers. It utilizes the tendency of betweenlayer representation transformations of ID data to be smoother than the corresponding transformations of OOD data. The smoothness of the transformation between layers l and $l + 1$ for an input

 x is quantified using the Frobenius norm of the Jacobian matrix for $l = 1, \ldots, L - 1$. This is given by:

$$
\phi_l(\boldsymbol{x}) = \|\boldsymbol{J}_l(\boldsymbol{h}_l)\|_F^2 = \sum_{i=1}^{d_{l+1}} \sum_{j=1}^{d_l} \left(\frac{\partial (f_{l+1})_i}{\partial (h_l)_j}\right)^2
$$
\n(15)

where $J_l(h_l)$ is the Jacobian matrix of the transformation from layer *l* to $l+1$, h_l is the representation at layer l, and $f_l : \mathbb{R}^{d_{l-1}} \to \mathbb{R}^{d_l}$ is the intermediate network layers, while f_L corresponds to the last layer, mapping to a vector of logits. To reduce computational complexity, in practice, BLOOD uses an unbiased estimator of the smoothness measure with r pairs of random vectors $v_l \sim \mathcal{N}(\mathbf{0}_n, \mathbf{I}_n)$ and $w_l \sim \mathcal{N}(\mathbf{0}_m, \mathbf{I}_m)$:

$$
\hat{\phi}_l(\boldsymbol{x}) = \frac{1}{r} \sum_{i=1}^r \left(\boldsymbol{w}_{l,i}^\top \boldsymbol{J}_l(\boldsymbol{h}_l) \boldsymbol{v}_{l,i} \right)^2 \qquad (16)
$$

The final BLOOD score for an input x can be computed as either the average smoothness score across all layers:

BLOOD_M =
$$
\frac{1}{L-1} \sum_{l=1}^{L-1} \hat{\phi}_l(x)
$$
 (17)

or the smoothness score at the last layer:

$$
\text{BLOOD}_L = \hat{\phi}_{L-1}(\boldsymbol{x}) \tag{18}
$$

We follow Jelenić et al. (2023) to use $BLOOD_L$ as the uncertainty score of an instance x for its higher performance. Finally, the OOD score is defined as:

$$
O(\mathbf{x}; E') = -\text{BLOOD}_L \tag{19}
$$

Note that this method is also applicable only when using fine-tuned encoder E' .

Mahalanobis The Mahalanobis distance detector proposed by [Lee et al.](#page-9-6) [\(2018\)](#page-9-6) is a widely used OOD detection method that calculates the OOD score $O(x; E)$ based on the distance of a test sample to the nearest ID class in the embedding space determined by M . It can be formulated as:

$$
O(\mathbf{x}; E) = \min_{c \in \{1, ..., C\}} (E(\mathbf{x}) - \boldsymbol{\mu}_c)^\top
$$

$$
\Sigma^{-1} (E(\mathbf{x}) - \boldsymbol{\mu}_c) \quad (20)
$$

where μ_c is the mean of all of the representations of the instances in class c and Σ is the covariance matrix. μ_c and Σ can be estimated by:

$$
\hat{\mu_c} = \frac{1}{N_c} \sum_{\boldsymbol{x} \in D_{IN}^c} E(\boldsymbol{x})
$$
\n(21)

$$
\hat{\Sigma} = \frac{1}{N} \sum_{c \in \{1, \dots, C\}} \sum_{\boldsymbol{x} \in D_{IN}^c} (22)
$$
\n
$$
(E(\boldsymbol{x}) - \boldsymbol{\mu}_c) (E(\boldsymbol{x}) - \boldsymbol{\mu}_c)^\top
$$

where $D_{IN}^c = \{ x \mid (x, y) \in D_{IN}, y = c \}$ represents for the training data belonging to the class c, N denotes the size of the training set, and N_c is the number of training data belonging to the class c.

 k -Nearest Neighbors (KNN) [Sun et al.](#page-10-3) [\(2022\)](#page-10-3) investigate the effectiveness of using nonparametric nearest-neighbor distances for OOD detection on visual OOD detection benchmarks. We applied this approach to text data, where $O(x; E)$ represents the distance from the test sample to its k-th nearest ID training sample in the normalized feature space. In our experiments, we set $k = 1$.

kMeans & C-kMeans We also compare our approaches to the standard kMeans algorithm and its class-wise variant, C-kMeans, similar to the C-NNK-Means. In both cases, we use the reconstruction error as the OOD score $O(x; E)$. The number of clusters is a hyper-parameter, and their selection will be discussed in [Appendix F.](#page-15-0)

D Implementation Details

Fine-tuning We fine-tuned the PLM for classification on the ID dataset and used the all-distilroberta-v1 checkpoint from HuggingFace. In all cases, we used mean-pooling on token representations from the penultimate layer to generate sentence-level representations. We used 5 different random seeds and reported the average results to limit the effect of randomness for each setting. All models were optimized with Cross Entropy Loss and AdamW [\(Loshchilov and Hutter,](#page-10-21) [2017\)](#page-10-21) as the optimizer, using a weight decay rate of 0.01 and a learning rate of 1×10^{-5} , with a linear learning rate decay. We used a batch size of 4 and fine-tuned the model for 5 epochs.

OOD Detection After extracting embeddings, we ran our baselines and proposed methods on a single NVIDIA Tesla V100 GPU to ensure consistent measurement of inference time. We tuned hyper-parameters based on the validation set and reported the final results on the test set of each dataset. [Appendix F](#page-15-0) provides more details of our hyper-parameter tuning.

E Evaluation Metrics

Here, we introduce 3 standard metrics for OOD detection and the Inference Time in seconds we used to compare the complexity:

AUROC The Area Under the Receiver Operating Characteristic Curve, plots the True Positive Rate (TPR) against the False Positive Rate (FPR) at various thresholds. A higher AUROC value indicates better performance.

AUPR The Area Under the Precision-Recall Curve, evaluates the model's precision and recall by plotting precision against recall for different thresholds. A higher AUPR value indicates better identification of OOD samples while maintaining high precision.

FPR@95 The False Positive Rate at 95% True Positive Rate, measures the FPR when the TPR is fixed at 95%. A lower FPR@95 value indicates fewer ID samples being misclassified as OOD, signifying a more reliable OOD detection model.

Inference Time It serves as an additional metric to account for the complexity of the OOD detection methods. We measured the time taken to obtain the OOD score of a given query q after extracting its representation from a PLM. Note that we do not report this for MSP, Energy, and D2U, as their inference involves minimally processing the logits, and so they have negligible inference time. We also do not report this for BLOOD since its inference process is significantly affected by the batch size. Additionally, BLOOD requires representations extracted from every layer of the model. So, despite doing limited processing after embeddings have been extracted, in practice, the complexity of this method is much higher than that of other classifier-based ones.

We provide the results of AUROC and Inference Time in Section [6,](#page-5-3) and AUPR and FPR@95 results in [Appendix A.](#page-11-3)

F Hyper-parameter Tuning

KMeans, NNK-Means, and EC-NNK-Means select the number of dictionary atoms from {500, 1000, 2000, 4000}. For the class-wise versions, C-kMeans, C-NNK-Means, and C-EC-NNK-Means, due to the smaller size of each class compared to the overall dataset, the selection range is {50, 150, 250, 350} instead. Additionally, for EC-NNK-Means and C-EC-NNK-Means, we also need to choose Entropy Constraint hyper-parameter λ from {50, 150, 250, 350}. We tuned the hyperparameters on the validation set of each dataset, selecting the optimal hyper-parameters based on AUROC for each dataset (and each known classes ratio), and obtained the final results on the test set. We applied the same hyper-parameter tuning process for the Pre-trained Embedding setting and the Margin-based Contrastive Loss Embedding setting. Detailed hyper-parameter choices for each setting can be found in [Table 9](#page-16-0) and [Table 10.](#page-16-1)

		20 Newsgroups				Banking77		AG News		CLINC150
	% ID Classes \rightarrow	25%	50%	75%	25%	50%	75%	50%	75%	
∽	kMeans C-kMeans NNK-Means C-NNK-Means EC-NNK-Means C-EC-NNK-Means	1000 250 2000 350 (2000, 0.03) (350, 0.01)	500 50 2000 350 (2000, 0.03) (350, 0.07)	500 50 4000 350 (2000, 0.03) (350, 0.03)	2000 32 1000 32 (2000, 0.03) (32, 0.01)	4000 32 2000 32 (4000, 0.01) (32, 0.01)	1000 32 4000 32 (2000, 0.03) (32, 0.01)	4000 350 4000 350 (4000, 0.03) (50, 0.07)	500 50 4000 50 (500, 0.01) (150, 0.07)	2000 50 2000 25 (2000, 0.05) (50, 0.01)

Table 9: Hyper-parameter settings for different methods with Cross Entropy Loss Embeddings on 4 datasets. This is used for our main results in Section [6.](#page-5-3) For EC-NNK-Means and C-EC-NNK-Means, the hyper-parameters are in the format of (M, λ) where M is the initial number of dictionary atoms and λ is the hyper-parameter that controls the influence of entropy-constrained term, while others are only using M .

Table 10: Hyper-parameter settings for different methods with Pre-trained Embeddings and Margin-based Contrastive Loss Embeddings on 20 Newsgroup. This is used for our additional analysis in Section [6](#page-5-3) to show the competitive performance of our methods with different embeddings. For EC-NNK-Means and C-EC-NNK-Means, the hyper-parameters are in the format of (M, λ) where M is the initial number of dictionary atoms and λ is the hyper-parameter that controls the influence of entropy-constrained term, while others are only using M.

Table 11: Example of data from an ID cluster from the visualization in [Figure 3,](#page-6-2) with identical label colors. Last column represents the NNK-Means Error, as presented in [\(9\)](#page-4-4). Bolded text is edited from the original to preserve anonymity.