EFFICIENT AND ROBUST NEURAL COMBINATORIAL OPTIMIZATION VIA WASSERSTEIN-BASED CORESETS

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Abstract

Combinatorial optimization (CO) is a fundamental tool in many fields. Many neural combinatorial optimization (NCO) methods have been proposed to solve CO problems. However, existing NCO methods typically require significant computational and storage resources, and face challenges in maintaining robustness to distribution shifts between training and test data. To address these issues, we model CO instances into probability measures, and introduce Wasserstein-based metrics to quantify the difference between CO instances. We then leverage a popular data compression technique, *coreset*, to construct a small-size proxy for the original large dataset. However, the time complexity of constructing a coreset is linearly dependent on the size of the dataset. Consequently, it becomes challenging when datasets are particularly large. Further, we accelerate the coreset construction by adapting it to the merge-and-reduce framework, enabling parallel computing. Additionally, we prove that our coreset is a good representation in theory. Subsequently, to speed up the training process for existing NCO methods, we propose an efficient training framework based on the coreset technique. We train the model on a small-size coreset rather than on the full dataset, and thus save substantial computational and storage resources. Inspired by hierarchical Gonzalez's algorithm, our coreset method is designed to capture the diversity of the dataset, which consequently improves robustness to distribution shifts. Finally, experimental results demonstrate that our training framework not only enhances robustness to distribution shifts but also achieves better performance with reduced resource requirements.

1 INTRODUCTION

Combinatorial optimization (CO) is a fundamental tool in many fields such as transportation (Contardo et al., 2012; Veres & Moussa, 2019), logistics (Laterre et al., 2018) and manufacturing (Froger et al., 2016; Dolgui et al., 2019; Liu et al., 2017). Numerous traditional exact (David Applegate, 2006; Optimization, 2020) or heuristic solvers (Croes, 1958; Helsgaun, 2017; Lamm et al., 2016) have been designed by experts to solve these problems. However, the real-world CO problems are widespread and diverse, and may even undergo rapid changes over time. Moreover, even for a fixed CO problem, human experts may be hindered by limited domain knowledge and computational difficulty (many of these CO problems are NP-hard). As a result, in many situations, it can be impractical to rely solely on hand-crafted methods developed by experts.

To address these challenges, numerous *Neural Combinatorial Optimization* (NCO) methods have been proposed, such as constructive heuristics methods (Khalil et al., 2017; Kool et al., 2018; Kwon et al., 2020; Hottung et al., 2020; Kim et al., 2022; Joshi et al., 2019; Fu et al., 2021; Geisler et al., 2021; Qiu et al., 2022; Sun & Yang, 2023; Luo et al., 2023; Vinyals et al., 2015; Bello et al., 2016; Nazari et al., 2018; Deudon et al., 2018; Xin et al., 2020; 2021; Kwon et al., 2021; Kim

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et al., 2021; Cheng et al., 2023; Drakulic et al., 2023) and improvement heuristics methods (Li et al., 2018; d O Costa et al., 2020; Wu et al., 2021; Chen & Tian, 2019; Li et al., 2023; Chen & Tian, 2019; Hottung et al., 2020; Joshi & Anand, 2022; Joshi et al., 2019). These methods learn heuristic solution strategies in a data-driven manner, thus dispensing with laborious manual design and expert knowledge; moreover, compared with traditional CO solvers, NCO methods can benefit from accelerated inference speeds by utilizing modern GPU devices.

Despite their advantages, these methods often require large training datasets, which demands substantial storage space and computational resources. Additionally, when training data and test data come from different distributions, enhancing the robustness to distribution shift (Liang et al., 2023; Sun et al., 2020) is also a challenge for existing NCO models. Therefore, training a competitive model with limited resources while ensuring its robustness to distribution shifts is an important and worthy problem to address.

To address these issues, we consider constructing a good representation, *coreset* (Ros & Guillaume, 2020), for the original huge dataset. Coreset is a popular data compression technique, which can accelerate the training process by reducing dataset size while preserving the value. Roughly speaking, coreset is a small-size proxy of the original dataset Q with respect to an objective; the value of the objective evaluated on coreset can closely approximate the value evaluated on Q. Therefore, we can replace Q by coreset in the training phase, and thus save the storage space and computational resources significantly. Furthermore, our coreset method is inspired by hierarchical Gonzalez's algorithm (Gonzalez, 1985), and thus can capture the diversity of the dataset. Consequently, benefiting from its diversity, the model based on our coreset method shows robustness to distribution shift.

The intuition behind our coreset technique can be likened to preparing for an exam. Training neural networks is similar to practicing exercises for an exam. While the number of available exercises (i.e., data) might be vast, we cannot be trained for all the exercises with limited time and energy (i.e., storage and computational resources). Fortunately, the whole exercises are redundant; to get a high score, doing all the exercises is unnecessary, and we only need to cover all categories of exercises. Based on the above intuitions, we need a small-size representation (i.e., coreset) for the whole exercises. To this end, three key steps are required: i) exploring a proper metric to quantify the difference between CO instances; ii) constructing a coreset for the original dataset; iii) designing an efficient training framework based on the coreset technique for existing NCO models.

Many CO problems, such as the Traveling Salesperson Problem (TSP) and Maximum Independent Set (MIS), can inherently induce a graph structure. By employing graph embedding techniques, we can map these graph structures into a set of points in Euclidean space. Thus, we model CO instances as probability measures (in section 3.1). Wasserstein distance is commonly used to quantify the difference between probability measures. However, solutions to CO problems such as TSP remain invariant under rigid transformations such as translation, rotation, and reflection. In other words, a CO instance can generate multiple variants through these transformations; but they are inherently the same instance. Therefore, the distance between such instances should be zero. To capture this property, we introduce the Wasserstein distance under **r**igid transformations (RWD) to measure the difference between two CO instances.

Our contributions:

- First, we model CO instances as probability measures, and introduce RWD to quantify the difference between two given CO instances.
- Then, based on RWD, we design a coreset algorithm to effectively compress data for training acceleration; it saves substantial computational and storage resources. However, the time required to construct the coreset increases linearly with the size of the dataset, making it computationally expensive for extremely large datasets.
- To further accelerate coreset construction, we adapt our coreset method to merge-andreduce framework, enabling parallel computation. Moreover, we demonstrate that our coreset is a good representation theoretically.
- Next, based on our coreset method, we propose an efficient training framework for accelerating the existing NCO training process. More specifically, we replace the original dataset with our coreset to accelerate the training process; in the inference phase, test instances are

aligned along our tree (i.e., \mathcal{T} from Algorithm 1 or 2) before predicting their labels using the trained model.

• The experimental results show that our training framework exhibits better performance and enhanced robustness to distribution shifts.

1.1 OTHER RELATED WORKS

Here, we introduce several techniques that will be involved later.

Graph embedding technique represents the nodes and edges of a graph in Euclidean space. The edge information is encoded within the Euclidean distances between points, reducing the need to handle complex graph structures directly, as point-to-point information suffices. Moreover, it transforms the discrete graph into continuous Euclidean coordinates, which allows many techniques in Euclidean space to be used. Here are some widely used graph embedding methods. Laplacian Eigenmaps (Belkin & Niyogi, 2001) embed graph data into a low-dimensional Euclidean space while preserving local neighborhood relationships. Multidimensional Scaling (MDS) (Borg & Groenen, 2007) focuses on preserving pairwise distances between nodes in the graph. Isomap (Tenenbaum et al., 2000) extends MDS by incorporating geodesic distances along the manifold, making it especially useful for graphs with inherent nonlinear structures.

Hierarchical Gonzalez's algorithm (Murtagh & Contreras, 2012) is a variant of Gonzalez's *k*-center algorithm for addressing hierarchical clustering problem. In this approach, clusters are recursively divided at different levels of granularity, yielding a tree structure for efficient querying. This algorithm prioritizes selecting new center points that are far apart from the previously chosen ones. This strategy leads to clusters well-spread across the data, effectively capturing the diversity of the dataset. This method is commonly used for summarizing large datasets. However, its time complexity exhibits a linear dependence on the size of the dataset, making it potentially time-consuming for extremely large datasets. To mitigate this issue, we integrate merge-and-reduce (Bentley & Saxe, 1980; Har-Peled & Mazumdar, 2004) technique to construct our coreset in Algorithm 2.

2 PRELIMINARIES

Notations We define $[n] := \{1, ..., n\}$ and denote the vector of ones by 1. The ℓ_2 -norm is denoted by $\|\cdot\|$, and |A| denotes the size of set A. Let \mathbb{R}_+ be the set of non-negative real numbers. Let $\mathcal{P}(\mathbb{R}^d)$ be the probability measure space on Euclidean space \mathbb{R}^d . Matrices are denoted by bold capital letters, such as C; C_{ij} is its element in the *i*-th row and *j*-th column. Similarly, we denote vectors by bold lowercase letters, such as $\mathbf{a} := (a_1, \ldots, a_n)^T \in \mathbb{R}^n$; a_i is its *i*-th element.

Wasserstein distance (Peyré et al., 2017) is skilled at capturing the geometric structures of CO problems, but it is sensitive to rigid transformations. To obtain the invariance property under rigid transformations, we consider the following conception: *Wasserstein distance under rigid transformation* (RWD).

Definition 2.1 (RWD). Let $\mu = \sum_{i=1}^{n} a_i \delta_{x_i}, \nu = \sum_{j=1}^{n} b_j \delta_{y_j} \in \mathcal{P}(\mathbb{R}^d)$, where $\mathbf{a}, \mathbf{b} \in \mathbb{R}^n_+$ are their weight vectors and $\{x_i\}_{i \in [n]}, \{y_j\}_{j \in [n]} \subset \mathbb{R}^d$ are their locations. Then, the Wasserstein distance under rigid transformation between μ and ν is

$$\mathcal{W}(\mu,\nu) := \left(\min_{\mathbf{P}\in\Pi(\mathbf{a},\mathbf{b}), e\in E(d)} \sum_{i=1}^{n} \sum_{j=1}^{n} P_{ij} \|x_i - e(y_j)\|^2 \right)^{1/2}$$

where $\Pi(\mathbf{a}, \mathbf{b}) := \{ \mathbf{P} \in \mathbb{R}^{n \times n}_+ | \mathbf{P}\mathbf{1} = \mathbf{a}, \mathbf{P}^T\mathbf{1} = \mathbf{b} \}$ is the coupling set, E(d) is the euclidean group on \mathbb{R}^d , and $e : \mathbb{R}^d \to \mathbb{R}^d$ is the rigid transformation.

Remark 2.2. *i)* If we fix e as identity transformation, then RWD is degenerated as the Wasserstein distance; Wasserstein distance is a metric on $\mathcal{P}(\mathbb{R}^d)$. *ii)* RWD is a (semi-)metric¹ on $\mathcal{P}(\mathbb{R}^d)$; more specifically, $(\mathcal{P}(\mathbb{R}^d), W)$ is a metric space.

¹For simplicity, we do not distinguish between metric and semi-metric.

Next, we formally define our coreset technique. Let

$$\ell: \mathcal{Q} \times \Theta \to \mathbb{R}_+, \quad (\mu, \theta) \mapsto \ell(\mu, \theta) \tag{1}$$

be a loss function, where $\theta \in \Theta$ is the model parameter and $\mu \in Q$ denotes a CO instance. For any weighted set $A \subset Q$ with weight function w_A , we define $\ell(A, \theta) := \sum_{\mu \in A} w_A(\mu) \cdot \ell(\mu, \theta)$.

Definition 2.3 (Coreset). Let $0 < \epsilon < 1$ and ℓ be a loss function. Let $\mathcal{Q} \subset \mathcal{P}(\mathbb{R}^d)$ be a set of measures with weight function $w_{\mathcal{Q}} : \mathcal{P}(\mathbb{R}^d) \to \mathbb{R}_+$. Let $\sum_{\mu \in \mathcal{Q}} w_{\mathcal{Q}}(\mu) = 1$. Then, a weighted set S with weight function w_S is an ϵ -coreset of \mathcal{Q} if

$$\ell(\mathcal{S},\theta) \in (1\pm\epsilon) \cdot \ell(\mathcal{Q},\theta) \quad \text{for all } \theta \in \Theta.$$
 (2)

Then, we introduce some basic properties that will be used later. *Doubling dimension* (Chan et al., 2016) can describe the growth rate of the dataset with respect to some metric dist. Formally, the doubling dimension of metric space (Q, dist) is the smallest positive integer ddim such that every ball in (Q, dist) can be covered by 2^{ddim} balls of half the radius. For example, the doubling dimension of the Euclidean space \mathbb{R}^d is $\Theta(d)$.

The Lipschitz constant of a function describes how fast it can change. The loss function is *L*-Lipschitz continuous with respect to dist on Q, if $|\ell(\mu_1, \theta) - \ell(\mu_2, \theta)| \le L \cdot \operatorname{dist}(\mu_1, \mu_2)$ holds for all $\mu_1, \mu_2 \in Q, \theta \in \Theta$.

3 OUR METHODS

This section introduces our methods. Section 3.1 introduces RWD to quantify the difference between two CO instances. Section 3.2 constructs a small-size coreset for accelerating the training process. Section 3.3 accelerates coreset construction process by using merge-and-reduce framework; moreover, we theoretically demonstrate that our coreset is a good representation. Finally, in Section 3.4, we present our efficient framework for existing NCO methods.

3.1 METRICS FOR CO INSTANCES

Many CO problems can induce graph structures. We first extract the graph structure induced by the CO instance and represent it by a graph metric space, where each point in this space reflects node-specific information, and edge relationships are captured through the corresponding shortest-path metric. We then apply graph embedding techniques (in Section 1.1) to map this graph metric space into Euclidean space, aiming to preserve inter-point distances closely. In this embedding, each node in the original graph is represented as a discrete point in Euclidean space, and edge information is encoded in Euclidean distances between these points. Ultimately, we represent the graph as a discrete set of points in Euclidean space. Henceforth, we focus on the point set data in Euclidean space.

Given two CO instances, we represent the nodes of their corresponding graph structure as $X = \{x_i\}_{i \in [n]}, Y = \{y_j\}_{j \in [n]} \subseteq \mathbb{R}^d$. Then, the CO instances are modeled as two probability measures $\mu = \sum_{i=1}^n a_i \delta_{x_i}$ and $\nu = \sum_{j=1}^n b_j \delta_{y_j}$ (with $a_i = b_j = \frac{1}{n}$ to represent equal node importance). Then, we can quantify the difference between μ and ν with metric RWD; that is, $\mathcal{W}(\mu, \nu)$, where the ground distance between $x \in X$ and $y \in Y$ is ||x - y|| as in Definition 2.1.

Remark 3.1. *i)* By graph embedding technique, the nodes and edges of a graph structure are described by the locations and their ground distances in Euclidean space. ii) These CO problems are usually invariant after imposing rigid transformations on their nodes, and RWD can capture this characteristic well. In essence, the complexity of data space is reduced under RWD metric. More specifically, we regard two CO instances as the same instance if their distance is zero. iii) The complete graph in \mathbb{R}^d can be represented directly by a point set, without the need for graph embedding techniques. iv) If we fixed the outer iteration number and the dimension d of data space as constants, our RWD can be solved within $\tilde{O}(n^2)$ time by using the heuristic method in Algorithm 3.

3.2 CORESET

Intuitively, our coreset aims to cover all the data by using relatively fewer and smaller balls, where all the balls have the same radius. This strategy can be likened to preparing for an exam, where

we need to cover as many categories of exercises as possible with limited time and energy. For simplicity, we take RWD as an example to illustrate our methods. The metric RWD can also be replaced by Wasserstein distance, or some other proper metrics on $\mathcal{P}(\mathbb{R}^d)$.



Figure 1: Grow nodes from root v_0 with ddim = 1. Figure 2: Accelerate the coreset construc-The red dots denote probability measures in dataset tion by using merge-and-reduce framework. Q, and the solid red dot is the center of the ball v.ball.

Coreset construction Our algorithm is inspired by the hierarchical structure in (Ding et al., 2021; Krauthgamer & Lee, 2004; Har-Peled & Mendel, 2005; Beygelzimer et al., 2006). Given a set Q of probability measures, we aim at clustering similar measures into small balls of radius r, and take the cluster centers to form our coreset. The coreset is finally reserved in the tree \mathcal{T} as shown in Figure 1.

To construct such a coreset, we first initialize an empty tree \mathcal{T} , and set its root node as v_0 . The root node has only one attribute buffer, and is initialized as v_0 .buffer = Q. Our (non-root) node has four attributes: cluster, center, buffer and ball as shown in Figure 1. The nodes grow in an up-bottom manner recursively. Given a current node v, if v.buffer is an empty set, then v is a leaf node, and we stop adding children to it. If v.buffer is a nonempty set, we add $k = \min\{|v.\text{buffer}|, 2^{2 \cdot \text{ddim}}\}$ children node $\{v'_j\}_{j \in [k]}$ to the current node v; more specifically, we run Gonzalez's algorithm k rounds on v.buffer. By this, we obtain k cluster centers v'_i .center and their corresponding clusters v'_i .cluster. All the v'_i .cluster form a partition of v.buffer; each v'_i .cluster consists of points that are relatively close to its center v'_j .center. For each set v'_j .cluster, we partition it into two sets v'_i .ball, v'_i .buffer, where v'_i .ball is a RWD-based ball of radius r centered at measure v'_i .center; formally, we can formulate them as

$$v'_{i}.\mathsf{ball} = \left\{ \mu \in v'_{i}.\mathsf{cluster} \mid \mathcal{W}(\mu, v'_{i}.\mathsf{center}) \le r \right\}$$
(3)

and

$$v'_i$$
.buffer = v'_i .cluster - v'_i .ball. (4)

Finally, we obtain a tree \mathcal{T} . The coreset \mathcal{S} consists of all the center points v.center with weight |v.ball|. We show the coreset construction process in a more intuitive and comprehensible manner in Figure 1. The detailed descriptions are in Algorithm 1.

Time complexity and coreset size From (Ding et al., 2021), we know that the radius of the clusters will be halved after carrying out at most $2^{2 \cdot \text{ddim}}$ rounds of Gonzalez's algorithm. Let R be the radius of dataset \mathcal{Q} ; that is, $\mathcal{W}(\mu,\nu) \leq 2R$ for any $\mu,\nu \in \mathcal{Q}$. Thus, the height of the tree in Algorithm 1 is at most $\mathcal{O}(\log \frac{R}{r})$. Let T(n) be the time for computing the distance (i.e., RWD) between two measures, where n is the size of the locations of measures. Since constructing every layer takes $\mathcal{O}(2^{2 \cdot \text{ddim}}) \cdot |\mathcal{Q}|$ computations of RWD, the total time complexity for Algorithm 1 is $\mathcal{O}(2^{2 \cdot \text{ddim}}) \cdot |\mathcal{Q}| \cdot T(n) \cdot \log \frac{R}{r}$. Its time complexity increases linearly with the size of the dataset, making it computationally expensive for large datasets. The coreset is maintained in the tee \mathcal{T} . The tree has $\mathcal{O}((2^{2 \cdot \text{ddim}})^{\log \frac{R}{r}})$ nodes, thus the coreset size is $\mathcal{O}((\frac{R}{r})^{2 \cdot \text{ddim}})$.

Remark 3.2. *i)* The output of Algorithm 1 contains a tree \mathcal{T} and coreset \mathcal{S} . The \mathcal{S} is a representation of the original measure set Q, which is used for speeding up the training process for existing NCO methods; while the tree T is prepared for aligning CO instances at the inference phase. ii) It is often not necessary to know the exact value of the doubling dimension in advance. Typically, we

Algorithm 1 Algorithm for constructing coresets

Input: a set $Q := {\{\mu_i\}}_{i \in [N]} \subset \mathcal{P}(\mathbb{R}^d)$ of measures, doubling dimension ddim of Q, radius r

- 1: Initialize an empty tree \mathcal{T} , and set its root node as v_0 ;
- 2: Set v_0 .buffer = Q;

 \triangleright The root node v_0 only has an attribute buffer, and it is not associated with any node.

- 3: Construct the nodes of \mathcal{T} recursively as follows: $\triangleright v$ is the current node.
- 4: if v.buffer is \emptyset then
- 5: The current node v is a leaf node, and we stop adding children to it;
- 6: **else**
- 7: Set $k = \min\{|v.\mathsf{buffer}|, 2^{2 \cdot \mathsf{ddim}}\}$ and add k children node $\{v'_j\}_{j \in [k]}$ to the current node v;
- 8: Run Gonzalez's algorithm k rounds on v.buffer. For each children node v'_j , we set its attributions cluster, center, ball, buffer according to Equation (3) and Equation (4);

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9: end if
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10: Set S = \{v.center \mid v \text{ is a node of } \mathcal{T}\} and set the weight as w_S(\mu) = |v.ball|;
Output:\mathcal{T}, S
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begin by experimenting with relatively small values, as demonstrated in our study where we set the low doubling dimension as ddim = 1. In practice, even if we cannot rigorously prove that the data satisfies low doubling dimension assumption, this generally does not impact the effectiveness of our experimental results. iii) All the v.ball consist of a partition of Q.

3.3 ACCELERATE THE CORESET CONSTRUCTION PROCESS

In this subsection, we adapt our coreset method to merge-and-reduce framework (Bentley & Saxe, 1980; Har-Peled & Mazumdar, 2004; Wang et al., 2021); by this, we offer a technique to accelerate our coreset construction process by achieving parallel computing; moreover, it can also be used to tackle streaming data.

Algorithm 2 is a combination of our coreset method and the merge-and-reduce framework as shown in Figure 2. We first set $s = O((\frac{R}{r})^{2 \cdot \text{ddim}})$, $H = \log_{\frac{r}{s}} \frac{|Q|}{s}$ and $r' = \frac{r}{H}$. The height of the tree in Figure 2 is at most H, and it is generated in a bottom-up manner.

We perform reduce and merge procedures alternatively in each epoch. More specifically, *reduce* means data compression; that is, we run Algorithm 1 by taking $(\mathcal{Q}_i^h, \operatorname{ddim}, r')$ as input, and obtain the corresponding coreset \mathcal{S}_i^h ; *merge* means putting together the coresets \mathcal{S}_i^h ; that is, $\mathcal{Q}^{h+1} = \bigcup_{i \in [m_h]} \mathcal{S}_i^h$. After *H* epochs, we obtain the coreset $\mathcal{S} = \mathcal{Q}^{H+1}$ and its corresponding tree $\mathcal{T} = \mathcal{T}^{H+1}$.

Algorithm 2 Algorithm for accelerating the coreset construction process

Input: a set $Q := \{\mu_i\}_{i \in [N]} \subset \mathcal{P}(\mathbb{R}^d)$ of measures, doubling dimension ddim of Q, radius r1: Set $s = \mathcal{O}((\frac{R}{r})^{2 \cdot ddim}), H = \log_{\frac{\tau}{s}} \frac{|Q|}{s}$ and $r' = \frac{r}{H}, Q^0 = Q$; 2: for h = 1, ..., H do 3: \triangleright reduce procedure Partition Q^h as $Q^h = \bigsqcup_{i \in [m_h]} Q_i^h$ with $|Q_i^h| = \mathcal{O}(\tau)$ and $m_h = \lceil \frac{|Q^h|}{\tau} \rceil$; For every Q_i^h , we run Algorithm 1 by taking $(Q_i^h, ddim, r')$ as input, and output $(\mathcal{T}_i^h, \mathcal{S}_i^h)$; 4: \triangleright merge procedure 5: $Q^{h+1} = \bigcup_{i \in [m_h]} \mathcal{S}_i^h$; 6: end for 7: Set $S = Q^{H+1}$ and $\mathcal{T} = \mathcal{T}^{H+1}$; Output: \mathcal{T}, S

Time complexity Given that the input size of Algorithm 1 is $\mathcal{O}(\tau)$, and its output size is *s*. Then, the tree induced by the merge-and-reduce framework has at most $H = \log_{\frac{\tau}{2}} \frac{|\mathcal{Q}|}{s}$ layers.

Each layer of Algorithm 2 performs multiple computations of Algorithm 1 in parallel. Hence, the time complexity of per layer is $\tilde{O}(2^{2 \cdot \text{ddim}} \cdot \tau \cdot T(n)) \cdot \log \frac{R}{r}$. Consequently, the overall time complexity of Algorithm 2 is $\tilde{O}(2^{2 \cdot \text{ddim}} \cdot \tau \cdot T(n) \cdot \log \frac{R}{r} \cdot \log_{\frac{T}{s}} \frac{|\mathcal{Q}|}{s})$, which is independent on the dataset size.

Communication complexity Our coreset is a subset of the original dataset, allowing us to transmit only the indexes of the CO instance items rather than the data items themselves. This significantly reduces transmission costs. As a result, the additional transfer complexity introduced by our merge-and-reduce framework in Algorithm 2 is, in practice, minimal and unlikely to pose a substantial overhead.

Moreover, the coreset size of Algorithm 2 remains consistent with that in Algorithm 1, it is sufficient to retain only the tree structure of the final layer in practice.

A good representation Next, we show that the coreset S is a good representation of the original huge dataset Q in the following theorem.

Theorem 3.3. Assume ℓ is L-Lipschitz continuous on $(\mathcal{Q}, \mathsf{RWD})$ and there exists $\gamma \in \mathbb{R}_+$ such that $\ell(\mathcal{Q}, \theta) \geq \gamma$ for all $\theta \in \Theta$. Let ddim be the doubling dimension of \mathcal{Q} with respect to RWD , and R be the radius of \mathcal{Q} . Then, by setting $r = \frac{\epsilon \gamma}{L}$, both Algorithm 1 and Algorithm 2 can generate an $\mathcal{O}((\frac{R}{r})^{2 \cdot \mathsf{ddim}})$ size ϵ -coreset S for the dataset \mathcal{Q} ; that is, for every θ , it holds that $\ell(S, \theta) \in (1 \pm \epsilon) \cdot \ell(\mathcal{Q}, \theta)$.

It shows that for every parameter $\theta \in \Theta$, the value of the loss function ℓ evaluated on small-size coreset S can approximate the value on the original dataset Q within $\mathcal{O}(\epsilon)$ relative error. Therefore, Theorem 3.3 demonstrates that our small-size coreset S is a good representation for the original huge dataset Q with respect to the objective ℓ .

Proof. Due to limited space, we only give the proof sketch here. More details are in Appendix. First, we prove that by taking (Q, ddim, r) as input, the coreset constructed by Algorithm 2 can cover the dataset Q by small balls of radius r by using mathematical induction. Then, we obtain that the value difference of the loss function between the data itself and its representation is small by Lipschitz continuous property. Third, by setting some parameters properly, we turn the additive error into a relative error and obtain an ϵ -coreset S.

3.4 AN EFFICIENT FRAMEWORK

Here, we introduce an efficient framework to train a comparative model by using limited resources for existing NCO methods. We first feed the original dataset Q into Algorithm 1 or Algorithm 2, and obtain the coreset S and tree T. The original dataset Q is replaced by small-size coreset S in the training phase. Thus, it saves the storage and computing resources significantly in the training phase. The probability measure μ in our coreset S has its own weight $w_S(\mu)$, which helps it represent the original dataset well. However, to capture the diversity better, we usually regard these data as equally important; that is, to improve the robustness to distribution shifts in experiments, we set their weights as $w_S = \frac{1}{|S|}$.



Figure 3: An efficient framework for accelerating existing NCO methods. The Algorithm 1 can be replaced by Algorithm 2, and Algorithm 4 is in Appendix.

Meanwhile, in the inference phase, we first align the test instances μ along our tree T, which aims to find a rigid transformation such that

$$\min_{e,\nu \in S} W(e(\mu),\nu),\tag{5}$$

where $W(\cdot, \cdot)$ is the Wasserstein distance and $e(\mu) := \sum_{i \in [n]} a_i \delta_{e(x_i)}$ for any $\nu = \sum_{j \in [n]} b_j \delta_{y_j}$. We offer a heuristic method (i.e., Algorithm 4 in Appendix) for the alignment as described in Equation (5). Thanks to the tree structure maintained by \mathcal{T} , we can finish the alignment for a test instance within $\mathcal{O}(k \cdot \log(|\mathcal{S}|) \cdot T(n))$ time. This is particularly efficient in practice since k is usually small. Without this tree structure, we would potentially need to align the test instance with every training data, which could be significantly more time-consuming.

Remark 3.4. By combining our coreset technique in the training phase and the alignment process in the test phase, we essentially reduce the complexity of data. Intuitively, if the model can solve an instance well, then it can solve similar instances well under metric RWD. Furthermore, our framework can be applied to other problems that inherently involve a graph structure. This demonstrates its general applicability across various domains where graph-based analysis is pertinent.

Remark 3.5. *i)* Our coreset only needs to be computed once, after which it can be used repeatedly to train different models and fine-tune parameters. ii) Even if the coreset computation is time-consuming, it is still valuable as it helps save storage space. iii) Our alignment process serves as an optional enhancement to improve performance rather than a mandatory step.

Sample size	ample size Method Test distribution		Gree	Greedy		+2-opt
Sample size	Method	Test distribution	Length (\downarrow)	Time (\downarrow)	Length (\downarrow)	Time (\downarrow)
		$\mathcal{N}(0,1)$	20.39	386	18.61	384
128000	Org	$\mathcal{N}(0, 4^2)$	76.41	374	67.39	388
		$\mathcal{U}(0,10)$	89.29	372	79.82	385
		$\mathcal{N}(0,1)$	22.34	378	18.92	387
	US	$\mathcal{N}(0, 4^2)$	101.95	379	69.28	395
		$\mathcal{U}(0,10)$	119.78	380	82.59	395
		$\mathcal{N}(0,1)$	22.21	372	18.87	379
4003	CS	$\mathcal{N}(0, 4^2)$	80.63	372	67.92	379
		$\mathcal{U}(0,10)$	94.73	373	80.64	377
		$\mathcal{N}(0,1)$	22.18	359	18.88	363
	CS-aligned	$\mathcal{N}(0,4^2)$	80.66	362	67.91	358
		$\mathcal{U}(0,10)$	94.94	361	80.53	360
	US	$\mathcal{N}(0,1)$	22.12	377	18.87	388
		$\mathcal{N}(0,4^2)$	83.17	377	68.13	378
		$\mathcal{U}(0,10)$	97.31	377	80.80	387
		$\mathcal{N}(0,1)$	21.79	366	18.84	383
8245	CS	$\mathcal{N}(0, 4^2)$	78.72	372	67.79	378
		$\mathcal{U}(0,10)$	92.99	374	80.35	377
		$\mathcal{N}(0,1)$	21.80	360	18.86	359
	CS-aligned	$\mathcal{N}(0,4^2)$	78.50	361	67.82	358
		$\mathcal{U}(0,10)$	93.04	355	80.42	361
		$\mathcal{N}(0,1)$	21.99	390	18.87	377
	US	$\mathcal{N}(0, 4^2)$	80.78	384	67.94	379
		$\mathcal{U}(0,10)$	95.01	369	80.60	379
		$\mathcal{N}(0,1)$	21.57	372	18.81	382
12951	CS	$\mathcal{N}(0,4^2)$	77.80	369	67.58	379
		$\mathcal{U}(0,10)$	92.01	378	80.23	375
		$\mathcal{N}(0,1)$	21.50	361	18.79	358
	CS-aligned	$\mathcal{N}(0,4^2)$	77.67	362	67.57	357
		$\mathcal{U}(0,10)$	92.01	358	80.21	359

Table 1: Comparison of uniform sampling and our coreset method using TSP100-2D- $\mathcal{N}(0, 1)$ as the training dataset on test data TSP100-2D from different distributions.

4 EXPERIMENTS WITH TSP

We take TSP100 as an example to show the advantages of our coreset method. All experiments are conducted on an NVIDIA L20 GPU. Due to limited space, further experiments (including TSP training on uniformly sampled data (Kool et al., 2018), the MIS problem (Ahn et al., 2020)) and Capacitated Vehicle Routing Problem (CVRP) (Nazari et al., 2018) are presented in the Appendix².

Table 2: Comparison of uniform sampling and our coreset method using TSP100-2D- $\mathcal{N}(0,1)$ as th	e
training dataset on test data of varying sizes. We fix the sample size as 12951.	

TSD size	Mathad	Test distribution	Gree	edy	Greedy-	+2-opt
I SF SIZE	Method	Test distribution	Length (\downarrow)	Time (\downarrow)	Length (\downarrow)	Time (\downarrow)
		$\mathcal{N}(0,1)$	33.69	109	27.14	112
	US	$\mathcal{N}(0, 4^2)$	125.99	108	96.70	112
		$\mathcal{U}(0,10)$	145.41	109	113.39	112
		$\mathcal{N}(0,1)$	30.75	107	26.69	110
TSP200	CS	$\mathcal{N}(0,4^2)$	110.48	109	94.84	111
		$\mathcal{U}(0,10)$	129.77	107	111.47	109
		$\mathcal{N}(0,1)$	30.77	77	26.68	79
	CS-aligned	$\mathcal{N}(0,4^2)$	110.99	78	94.59	79
		$\mathcal{U}(0,10)$	129.28	76	111.49	78
		$\mathcal{N}(0,1)$	59.81	1012	43.41	1020
	US	$\mathcal{N}(0,4^2)$	237.72	1012	154.28	1022
		$\mathcal{U}(0,10)$	263.66	1015	180.75	1022
	CS	$\mathcal{N}(0,1)$	49.11	1012	42.25	1016
TSP500		$\mathcal{N}(0,4^2)$	178.56	1010	149.50	1016
		$\mathcal{U}(0,10)$	208.36	1011	174.93	1016
		$\mathcal{N}(0,1)$	49.38	680	42.26	683
	CS-aligned	$\mathcal{N}(0,4^2)$	178.88	679	149.63	682
		$\mathcal{U}(0,10)$	208.77	678	175.03	682
		$\mathcal{N}(0,1)$	94.71	2823	61.72	2848
	US	$\mathcal{N}(0,4^2)$	382.77	4224	219.16	2847
		$\mathcal{U}(0,10)$	426.61	4215	255.95	4254
		$\mathcal{N}(0,1)$	69.76	2823	59.59	2833
TSP1000	CS	$\mathcal{N}(0,4^2)$	252.92	4224	210.71	2832
		$\mathcal{U}(0,10)$	299.80	4215	246.57	4234
		$\mathcal{N}(0,1)$	69.96	2825	59.57	2832
	CS-aligned	$\mathcal{N}(0,4^2)$	253.63	2821	210.81	2830
		$\mathcal{U}(0,10)$	300.03	2812	246.61	2827

Dataset We apply our method on TSP100-2D/3D Euclidean instances. The labels of TSP100-2D instances are obtained by using the LKH-3 heuristic solver (Helsgaun, 2017); each coordinate of the nodes in a TSP instance is generated by x%10, where x is randomly sampled either from a normal distribution $\mathcal{N}(0, \sigma^2)$ or uniform sampling $\mathcal{U}(0, 10)$. Our training data, TSP100-2D- $\mathcal{N}(0, 1)$ and TSP100-3D- $\mathcal{N}(0, 1)$, consists of 125,000 instances generated by the normal distribution $\mathcal{N}(0, 1)$ and 3000 instances by the uniform distribution $\mathcal{U}(0, 10)$. Indeed, the distribution of the training dataset is very close to the normal distribution $\mathcal{N}(0, 1)$. Hence, we regard the test data sampled from distribution $\mathcal{N}(0, 1)$ as having no distribution shift. The test data are sampled from a single distribution, either a normal distribution or uniform distribution. Specifically, we sample 1280 test data items for TSP100, and 128 test data items for other cases. Obviously, the uniform distribution has the highest entropy and thus the highest diversity. For these Gaussian distributions, the larger the variance, the larger the diversity.

As for the TSP100-3D dataset, we can directly extend the 2D instances to 3D instances by appending a third coordinate with a value of zero. Let $X = \{x_i\}_{i \in [n]} \subset \mathbb{R}^3$ are the nodes of an instance. We apply random rotation transformation e on X; that is, $e(X) := \{e(x_i)\}_{i \in [n]}$.

²Code can be found at Coreset2025.

Setting We use the DIFUSCO (Sun & Yang, 2023) as our NCO solver. The detailed parameter settings are in Appendix. To quantify the performance of different methods, we use two criteria: the average tour length (Length) and the total runtime (Time). The term "Greedy" refers to the greedy decoding method of DIFUSCO, and "2-opt" is a post-processing used to improve solutions. The terms US,CS and CS-aligned represent uniform sampling, our coreset without alignment, and our coreset with alignment, respectively. We take the model trained on the full dataset with 128000 data items as baseline.

Samula siza	Mathad	Test distribution	Greedy		Greedy+2-opt	
Sample size	Method		Length (\downarrow)	Time (\downarrow)	Length (\downarrow)	Time (\downarrow)
128000	Org	$\mathcal{N}(0,1)$	129.35	108	112.23	106
	US	$\mathcal{N}(0,1)$	190.79	109	115.87	108
4003	CS	$\mathcal{N}(0,1)$	153.08	107	113.56	108
	CS-aligned	$\mathcal{N}(0,1)$	152.70	103	113.71	105
	US	$\mathcal{N}(0,1)$	166.40	106	114.47	108
8245	CS	$\mathcal{N}(0,1)$	140.49	107	113.04	106
	CS-aligned	$\mathcal{N}(0,1)$	140.18	104	112.91	104
	US	$\mathcal{N}(0,1)$	162.19	107	114.31	107
12951	CS	$\mathcal{N}(0,1)$	133.63	106	112.45	105
	CS-aligned	$\mathcal{N}(0,1)$	133.14	103	112.52	110

Table 3: Comparison of uniform sampling and our coreset method using TSP100-2D- $\mathcal{N}(0, 1)$ as the training dataset on test data TSPLIB(Reinelt, 1991).

Results of TSP100-2D Tables 1 to 3 present the results on training dataset TSP100-2D- $\mathcal{N}(0, 1)$. The training datasets are generated by the uniform sampling and our coreset technique respectively. The results show that both our method and uniform sampling method perform better as the sample size increases. Meanwhile, as the sample size decreases, the advantage of our methods compared to uniform sampling becomes increasingly evident.

Moreover, Tables 1 and 2 demonstrate that our method is robust to distribution shift. Specifically, the training data is sampled from a normal distribution $\mathcal{N}(0, 1)$, while the test data are sampled from normal distributions $\mathcal{N}(0, 1)$, $\mathcal{N}(0, 4^2)$ and a uniform distribution $\mathcal{U}(0, 10)$. The test distributions $\mathcal{N}(0, 4^2)$ and $\mathcal{U}(0, 10)$ are significantly different from the training distribution $\mathcal{N}(0, 1)$, which represent substantial distribution shifts. The results in Tables 1 and 2 show that our method consistently outperforms the baselines, demonstrating its robustness to distribution shifts.

Furthermore, Table 2 shows that models trained on our coreset can generalize better to larger problem sizes such as TSP200, TSP500 and TSP1000. Moreover, Table 3 confirms that our method outperforms other approaches on the TSPLIB dataset.

Results of TSP100-3D Tables 8 to 10 show the results on training dataset TSP100-3D- $\mathcal{N}(0, 1)$. For the test data without distribution shift (i.e., $\mathcal{N}(0, 1)$), our method has comparable performance; for the test data occurring distribution shift (i.e., $\mathcal{N}(0, 4^2)$ and $\mathcal{U}(0, 10)$), our method has better performance. Moreover, our coreset with alignment version performs better in the TSP-3D case. From Tables 1 to 3, 9 and 10, alignment can perform better for higher dimension dataset (i.e., TSP-3D). Thus, the alignment version is promising for tackling high-dimensional data. (The details are in Appendix.)

5 CONCLUSION AND FUTURE WORK

In this paper, we introduce an efficient training framework for NCO problems based on our coreset method. More specifically, we replace the original huge dataset with our coreset during the training phase. In the test phase, we first align the test instances with the data in our coreset, and then feed them into existing NCO models. Our framework enables the development of comparable models with limited computational and storage resources; additionally, it exhibits robustness to distribution shifts. Moreover, in future work, we will extend our method to other situations that can induce graph structures.

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A OTHER PRELIMINARIES

Lemma A.1 (Generalized triangle inequalities(Makarychev et al., 2022)). *Given three points* a, b, c, *the following inequalities hold for any* $0 < t \le 1$:

•
$$\operatorname{dist}^{2}(a,b) \leq (1+t) \cdot \operatorname{dist}^{2}(a,c) + (1+\frac{1}{t}) \cdot \operatorname{dist}^{2}(b,c);$$

• $\left|\operatorname{dist}^2(a,c) - \operatorname{dist}^2(b,c)\right| \le t \cdot \operatorname{dist}^2(a,c) + \frac{6}{t} \cdot \operatorname{dist}^2(a,b).$

Definition A.2 (Wasserstein distance (Peyré et al., 2017)). Let $\mu = \sum_{i=1}^{n} a_i \delta_{x_i}, \nu = \sum_{j=1}^{n} b_j \delta_{y_j} \in \mathcal{P}(\mathbb{R}^d)$, where $\mathbf{a}, \mathbf{b} \in \mathbb{R}^n_+$ are their weights and $\{x_i\}_{i \in [n]}, \{y_j\}_{j \in [n]} \subset \mathbb{R}^d$ are their locations. Given a cost matrix $\mathbf{C} \in \mathbb{R}^{n \times n}_+$ with $C_{ij} = \|x_i - y_j\|^2$, the Wasserstein distance between μ and ν is

$$W(\mu,\nu) := \left(\min_{\mathbf{P}\in\Pi(\mathbf{a},\mathbf{b})} \langle \mathbf{P},\mathbf{C} \rangle\right)^{1/2}$$

where $\Pi(\mathbf{a}, \mathbf{b}) := \{ \mathbf{P} \in \mathbb{R}^{n \times n}_+ \mid \mathbf{P}\mathbf{1} = \mathbf{a}, \mathbf{P}^T\mathbf{1} = \mathbf{b} \}$ is the coupling set and $\mathbf{1}$ is the vector of ones.

A.1 OTHER RELATED WORKS

NCO The existing NCO methods can be categorized into two types: constructive heuristics (Khalil et al., 2017; Kool et al., 2018; Kwon et al., 2020; Hottung et al., 2020; Kim et al., 2022; Joshi et al., 2019; Fu et al., 2021; Geisler et al., 2021; Qiu et al., 2022; Sun & Yang, 2023; Luo et al., 2023; Vinyals et al., 2015; Bello et al., 2016; Nazari et al., 2018; Deudon et al., 2018; Xin et al., 2020; 2021; Kwon et al., 2021; Kim et al., 2021; Cheng et al., 2023; Drakulic et al., 2023) and improvement heuristics (Li et al., 2018; d O Costa et al., 2020; Wu et al., 2021; Chen & Tian, 2019; Li et al., 2020; Joshi & Anand, 2022; Joshi et al., 2019) methods; the former can be further divided into two subtypes: autoregressive methods and non-autoregressive methods. The autoregressive methods directly predict a heatmap. The improvement heuristic methods often work by iteratively improving a feasible initial solution.

Coreset Coreset is a popular data compression technique for clustering Chen (2009); Feldman & Langberg (2011); Braverman et al. (2022), regression Tukan et al. (2020) and optimization Huang et al. (2022); Wang et al. (2021). More relevant, Huang et al. Huang et al. (2021) proposed a sequential coreset for optimization problems with the Lipschitz smoothness property. Wang et al. (Wang et al., 2021) designed a coreset method for continuous-and-bounded learning (Shalev-Shwartz & Ben-David, 2014). However, these methods cannot offer an efficient method for aligning CO instances with training data in the inference phase.

Optimal transportation (OT) Discrete Wasserstein distance is a special case of OT, thus it can be computed by standard OT solvers. In recent years, a lot of algorithms have been proposed to solve OT problem. For example, interior point method can compute an ϵ_+ -approximation value for OT with $\tilde{\mathcal{O}}(n^3)$ time in practice (Peyré et al., 2017) or $\tilde{\mathcal{O}}(n^{2.5})$ in theory (Lee & Sidford, 2015). To obtain an ϵ_+ -approximation solution of OT, Sinkhorn algorithm takes $\tilde{\mathcal{O}}(n^2/\epsilon_+^2)$ time (Dvurechensky et al., 2018; Lin et al., 2019) by solving the entropic regularization OT (Cuturi, 2013); the accelerated version of Sinkhorn algorithm yields $\tilde{\mathcal{O}}(n^{2.5}/\epsilon_+)$ time (Guminov et al., 2020); especially, based on area-convexity and dual extrapolation, Jambulapati et al. (2019) achieved $\tilde{\mathcal{O}}(n^2/\epsilon_+)$ time complexity.

B OTHER ALGORITHMS

Algorithm for computing RWD We define two discrete probability measures

$$\alpha = \sum_{i=1}^{n} a_i \delta_{x_i}, \beta = \sum_{j=1}^{n} b_j \delta_{y_j} \in \mathcal{P}(\mathbb{R}^d),$$
(6)

where $\mathbf{a}, \mathbf{b} \in \mathbb{R}^n_+$ are their weights and $\{x_i\}_{i \in [n]}, \{y_j\}_{j \in [n]} \subseteq \mathbb{R}^d$ their locations; δ is the Dirac delta function. We denote the Wasserstein distance between α and β by $W(\alpha, \beta)$.

The aim of Algorithm 3 is to find a rigid transformation e such that $\mathcal{W}(\alpha,\beta) = W(e \circ \alpha,\beta)$. We initialize $\tilde{\alpha} = \alpha$. We solve $\mathcal{W}(\alpha,\beta)$ by updating the coupling **P** and rigid transformation e alternatively. Specifically, we **first** obtain coupling **P** by computing $W(\tilde{\alpha},\beta)$ according to the method in (Jambulapati et al., 2019). **Then**, we fix **P**, and compute $\arg \min_e \mathcal{W}_{\mathbf{P}}(e \circ \alpha, \beta)$.

We partition the rigid transformation e into translation transformation e_1 and orthogonal transformation e_2 ; that is, $e = e_2 \circ e_1$. The translation transformation can be updated as $e_1 = \sum_{i=1}^{n} \sum_{j=1}^{n} P_{ij}y_j - \sum_{i=1}^{n} \sum_{j=1}^{n} P_{ij}x_i$. For fixed e_1 , **P**, computing the optimal orthogonal transformation e_2 is an orthogonal Procrustes problem (Gower & Dijksterhuis, 2004). More specifically, we first obtain $\mathbf{M} = \sum_{ij} P_{ij}x_iy_j^T$, and apply singular value decomposition $\mathbf{M} = \mathbf{UDV}^T$; then, we have

$$e_2 = \mathbf{U}\mathbf{V}^T. \tag{7}$$

Algorithm 3 Algorithm for RWD

Input: α, β 1: $t = 0, \tilde{\alpha} = \alpha$; 2: for $t < T_{align}$ do 3: t = t + 1;4: \triangleright update coupling **P** Obtain coupling **P** by computing $W(\tilde{\alpha}, \beta)$ according to the method in (Jambulapati et al., 2019); \triangleright update translation transformation e_1 and orthogonal transformation e_2 5: Compute $e_1 = \sum_{i=1}^n \sum_{j=1}^n P_{ij} y_j - \sum_{i=1}^n \sum_{j=1}^n P_{ij} x_i;$ Compute $e_2 = \mathbf{U}\mathbf{V}^T$ according to Equation (7), and set $e = e_2 \circ e_1$; $\tilde{\alpha} = \sum_{i=1}^{n} a_i \delta_{(e \circ x_i)}.$ 6: end for Output:P, e

Time complexity of Algorithm 3 We compute the RWD by alternating between optimizing the coupling matrix and the rigid transformation, which is a heuristic method. We assuming that the point dimension d and the number of iterations T_{align} are constants. For computing the coupling matrix, we solve an OT problem within $\tilde{O}(n^2)$ time (Jambulapati et al., 2019). The rigid transformation is obtained by solving an orthogonal Procrustes problem, which has a time complexity of $O(n^2d + nd^2 + d^3)$. Thus, the overall complexity of this heuristic method remains $\tilde{O}(n^2)$.

Algorithm for alignment Here, we introduce our alignment algorithm. We first initialize the current node v as the root node v_0 . Then, we walk from the root node to a leaf node by selecting the most similar measure with the test instance μ .

Algorithm 4 Algorithm for alignment

Input: the tree \mathcal{T} , CO instance μ 1: $v = v_0 \triangleright$ The current node v is initialized as root node v_0 . 2: Initialize *target* by any measure in coreset S. \triangleright S is the corresponding coreset maintained in tree \mathcal{T} . 3: Select the child v' that is closest to μ under metric RWD. 4: Set v = v'. 5: if $\mathcal{W}(\mu, v'.\text{center}) \leq \mathcal{W}(\mu, target)$ then 6: target = v'.center;7: end if 8: If v is not a leaf, jump to Line 3. **Output:**target



Figure 4: The illustration for error.

C OMITED PROOFS

Proof of Theorem 3.3. The proof sketch is listed here. First, we prove that by taking (Q, ddim, r) as input, the coreset constructed by Algorithm 2 can cover the dataset Q by small balls of radius r by using mathematical induction. Then, we obtain that the value difference of the loss function between the data itself and its representation is small by Lipschitz continuous property. Third, by setting some parameters properly, we turn the additive error into a relative error, and obtain an ϵ -coreset S.

Given a probability measure $\mu \in Q$, we assume that its corresponding ball center in the *h*-th epoch is ν_h .

Claim C.1. $\mathcal{W}^2(\mu,\nu_h) \leq h^2 \cdot \frac{r^2}{H^2}$.

Proof. The error is grown in a manner illustrated in Figure 4. Next, we prove this claim by using mathematical induction.

Base Case: For the case h = 1, in the 1-st epoch, we have $\mathcal{W}^2(\mu, \nu_1) \leq \frac{r^2}{H^2}$.

Induction step: For the case h = m, in the *m*-th epoch, we assume that

$$\mathcal{W}^2(\mu,\nu_m) \le m^2 \cdot \frac{r^2}{H^2}.$$
(8)

Then, according to the generalized triangle inequalities in Lemma A.1, we have

$$\mathcal{W}^{2}(\mu,\nu_{m+1}) \leq (1+t) \cdot \mathcal{W}^{2}(\mu,\nu_{m}) + (1+\frac{1}{t}) \cdot \mathcal{W}^{2}(\nu_{m},\nu_{m+1}).$$
(9)

Since the radius of small ball in Figure 4 is at most $\frac{r}{H}$, we have $\mathcal{W}^2(\nu_m, \nu_{m+1}) < \frac{r^2}{H^2}$. By using the the induction hypothesis and setting $t = \frac{1}{m}$, we have

$$\mathcal{W}^2(\mu,\nu_{m+1}) \le (1+\frac{1}{m}) \cdot m^2 \cdot \frac{r^2}{H^2} + (1+m) \cdot \frac{r^2}{H^2} = (m+1)^2 \frac{r^2}{H^2}.$$
 (10)

Till now, we prove the case h = m + 1.

Let $\nu := \nu_H$ be the representation of μ in Algorithm 2. According to Claim C.1, we have $W^2(\mu, \nu) \leq r^2$.

Next, since loss function is L-Lipschitz continuous with respect to RWD on Q, we have

$$|\ell(\mu,\theta) - \ell(\nu,\theta)| \le L \cdot \mathcal{W}(\nu,\mu) \le L \cdot r.$$
(11)

$$|\ell(\mathcal{Q},\theta) - \ell(\mathcal{S},\theta)| \le \sum_{\mu \in \mathcal{Q}} w_{\mathcal{Q}}(\mu) \cdot |\ell(\mu,\theta) - \ell(\nu,\theta)| \le \sum_{\mu \in \mathcal{Q}} w_{\mathcal{Q}}(\mu) \cdot L \cdot r = L \cdot r,$$
(12)

where $\nu \in S$ is the corresponding representation of $\mu \in Q$, and we have $w_Q(Q) := \sum_{\mu} w_Q(\mu) = 1$ according to Definition 2.1.

Finally, we have $|\ell(\mathcal{Q}, \theta)| \geq \gamma$, by setting $r = \frac{\epsilon \gamma}{L}$, we obtain the coreset property

$$|\ell(\mathcal{Q},\theta) - \ell(\mathcal{S},\theta)| \le \epsilon \gamma \le \epsilon \cdot |\ell(\mathcal{Q},\theta)|.$$
(13)

D FULL EXPERIMENTS WITH TSP

We take TSP100 as an example to show the advantages of our coreset method. All experiments are conducted on an NVIDIA L20 GPU. We take the NCO Sun & Yang (2023) as our backbone solver, and set its training epoch as 20.

Dataset We apply our method on TSP100-2D/3D Euclidean instances. The labels of TSP100-2D instances are obtained by using the LKH-3 heuristic solver (Helsgaun, 2017); each coordinate of the nodes in a TSP instance is generated by x%10, where x is randomly sampled either from a normal distribution $\mathcal{N}(0, \sigma^2)$ or uniform sampling $\mathcal{U}(0, 10)$. Our training data, TSP100-2D- $\mathcal{N}(0, 1)$ and TSP100-3D- $\mathcal{N}(0, 1)$, consists of 125,000 instances generated by the normal distribution $\mathcal{N}(0, 1)$ and 3000 instances by the uniform distribution $\mathcal{U}(0, 10)$. While the training dataset TSP100-2D- $\mathcal{U}(0, 10)$ consists of 128000 instances generated by uniform distribution $\mathcal{U}(0, 10)$.

Indeed, the distributions of the training dataset TSP100-2D- $\mathcal{N}(0,1)$ and TSP100-3D- $\mathcal{N}(0,1)$ are very close to the normal distribution $\mathcal{N}(0,1)$. Hence, we regard the test data sampled from distribution $\mathcal{N}(0,1)$ as having no distribution shift. The test data are sampled from a single distribution, either a normal distribution or uniform distribution. Specifically, we sample 1280 test data for TSP100, and 128 test data for other cases. Obviously, the uniform distribution has the highest entropy and thus the highest diversity. For these Gaussian distributions, the larger the variance, the larger the diversity.

As for the TSP100-3D dataset, we can directly extend the 2D instances to 3D instances by appending a third coordinate with a value of zero. Let $X = \{x_i\}_{i \in [n]} \subset \mathbb{R}^3$ are the nodes of an instance. We apply random rotation transformation e on X; that is, $e(X) := \{e(x_i)\}_{i \in [n]}$. Intuitively, this TSP100-3D dataset has the same intrinsic complexity under metric RWD, which is the low doubling dimension in our assumption.

Method	Sample size	Labeling time	Coreset Time	Training Time	Total time
Org	128000	4709	-	28563	33272
	4003	147	-	1894	2041
US	8245	304	-	2862	3166
	12951	475	-	4014	4489
	4003	145	691	1731	2567
CS	8245	305	1086	2747	4138
	12951	474	1283	3751	5508

Table 4: Time statistics for different phases of training on TSP100-2D- $\mathcal{N}(0,1)$.

Setting We use the DIFUSCO (Sun & Yang, 2023) as our NCO solver. The model DIFUSCO with Greedy decoding solves TSP instances in an end-to-end manner. We set the learning rate as 0.0002 and the batch size as 64. The diffusion step is performed 50 times in inference phase. We use the cosine schedule described in (Sun & Yang, 2023). To quantify the performance of different methods, we use two criteria: the average tour length (Length) and the total runtime (Time). The term "Greedy" refers to the greedy decoding method of DIFUSCO, and "2-opt" is a post-processing used to improve solutions. The terms US,CS and CS-aligned represent uniform sampling, our coreset without alignment, and our coreset with alignment, respectively. We use the results on the full dataset with 128000 data as a baseline. In experiments, we first construct a coreset S, and then take |S| samples by uniform sampling as training datasets.

Results of TSP100-2D Tables 4 to 7 present the results on training dataset TSP100-2D- $\mathcal{N}(0, 1)$. The training datasets are generated by the uniform sampling and our coreset technique respectively.

G 1	M. 41 1	The station of the state of the	Gree	edy	Greedy-	+2-opt
Sample size	Method	lest distribution	Length (\downarrow)	Time (\downarrow)	Length (\downarrow)	Time (\downarrow)
		$\mathcal{N}(0,1)$	20.39	386	18.61	384
		$\Lambda(0, 2^2)$	42 41	381	37.47	387
128000	Org	N(0, 2)	76.41	374	67.39	388
120000	018	N(0, 1)	87.18	379	77.86	388
		$\frac{1}{2}(0, 0)$	89.29	372	79.82	385
		$\frac{N(0, 10)}{N(0, 1)}$	22.34	378	18.92	387
		$\mathcal{N}(0, 2^2)$	51 59	376	38.25	388
	US	$\mathcal{N}(0, 2^{2})$	101 95	379	69.28	395
		$\mathcal{N}(0, 8^2)$	118.83	379	80.38	402
		$\mathcal{U}(0,0)$	119.78	380	82.59	395
		$\frac{N(0, 10)}{N(0, 1)}$	22.21	372	18.87	379
		$\mathcal{N}(0, 2^2)$	44.94	379	37.80	378
4003	CS	$\mathcal{N}(0, 4^2)$	80.63	372	67.92	379
		$N(0, 8^2)$	92.63	367	78 47	378
		$\mathcal{U}(0,0)$	94.73	373	80.64	377
		$\frac{N(0, 10)}{N(0, 1)}$	22.18	359	18.88	363
		$\mathcal{N}(0,2^2)$	45.00	357	37.80	362
	CS-aligned	$\mathcal{N}(0, 2^{-})$	80.66	362	67.91	358
	8	$\mathcal{N}(0, 8^2)$	92.59	362	78.41	358
		$\mathcal{U}(0, 10)$	94 94	361	80.53	360
		$\frac{N(0, 10)}{N(0, 1)}$	22.12	377	18.87	388
		$\mathcal{N}(0, 2^2)$	45.59	381	37.86	389
	US	$\mathcal{N}(0, 4^2)$	83.17	377	68.13	378
		$\mathcal{N}(0, 8^2)$	95.16	380	78.81	385
		$\mathcal{U}(0, 10)$	97.31	377	80.80	387
		$\frac{N(0,10)}{N(0,1)}$	21.79	366	18.84	383
		$\mathcal{N}(0,2^2)$	43.72	373	37.73	378
8245	CS	$\mathcal{N}(0, 4^2)$	78.72	372	67.79	378
		$\mathcal{N}(0, 8^2)$	90.44	371	78.36	380
		$\mathcal{U}(0,10)$	92.99	374	80.35	377
		$\mathcal{N}(0,1)$	21.80	360	18.86	359
		$\mathcal{N}(0,2^2)$	43.77	354	37.73	356
	CS-aligned	$\mathcal{N}(0,4^2)$	78.50	361	67.82	358
	C	$\mathcal{N}(0, 8^2)$	90.54	350	78.32	359
		$\mathcal{U}(0,10)$	93.04	355	80.42	361
		$\mathcal{N}(0,1)$	21.99	390	18.87	377
		$\mathcal{N}(0, 2^2)$	44.77	376	37.81	384
	US	$\mathcal{N}(0, 4^2)$	80.78	384	67.94	379
		$\mathcal{N}(0, 8^2)$	93.16	373	78.52	381
		$\mathcal{U}(0, 10)$	95.01	369	80.60	379
		$\mathcal{N}(0,1)$	21.57	372	18.81	382
		$\mathcal{N}(0,2^2)$	43.14	371	37.66	388
12951	CS	$\mathcal{N}(0, 4^2)$	77.80	369	67.58	379
		$\mathcal{N}(0,8^2)$	89.63	371	78.18	408
		$\mathcal{U}(0,10)$	92.01	378	80.23	375
		$\mathcal{N}(0,1)$	21.50	361	18.79	358
		$\mathcal{N}(0,2^2)$	43.18	361	37.66	364
	CS-aligned	$\mathcal{N}(0,4^2)$	77.67	362	67.57	357
		$\mathcal{N}(0,8^2)$	89.60	357	78.18	361
		$\mathcal{U}(0, 10)$	92.01	358	80.21	359

Table 5:	Comparison of uniform sampling and our coreset method using TSP100-2D- $\mathcal{N}(0,1)$ as
the trainir	ng dataset on test data TSP100-2D from different distributions.

The results show that both our method and uniform sampling method perform better as the sample size increases. Meanwhile, as the sample size decreases, the advantage of our methods compared to uniform sampling becomes increasingly evident.

Moreover, Tables 5 and 6 demonstrate that our method is robust to distribution shift. Specifically, the training data is sampled from a normal distribution $\mathcal{N}(0,1)$, while the test data is sampled from normal distributions $\mathcal{N}(0,1)$, $\mathcal{N}(0,4^2)$ and a uniform distribution $\mathcal{U}(0,10)$. The test distributions $\mathcal{N}(0,4^2)$ and $\mathcal{U}(0,10)$ are significantly different from the training distribution $\mathcal{N}(0,1)$, which represent substantial distribution shifts. The results in Tables 5 and 6 show that our method consistently outperforms the baselines, demonstrating its robustness to distribution shifts.

Furthermore, Table 6 shows that models trained on our coreset can generalize better to larger problem sizes such as TSP200, TSP500 and TSP1000. Moreover, Table 7 confirms that our method outperforms other approaches on the TSPLIB dataset. Table 4 shows the time efficiency of our coreset method.

		The different disc	Gree	Greedy		Greedy+2-opt	
I SP size	Method	lest distribution	Length (\downarrow)	Time (\downarrow)	Length (\downarrow)	Time (\downarrow)	
		$\mathcal{N}(0,1)$	29.85	107	26.61	111	
		$\mathcal{N}(0, 2^2)$	60.87	109	53.38	110	
TSP-200	Org	$\mathcal{N}(0, 4^2)$	109.70	110	94.90	111	
	_	$\mathcal{N}(0, 8^2)$	123.39	110	108.41	110	
		$\mathcal{U}(0, 10)$	126.99	107	111.50	111	
		$\mathcal{N}(0,1)$	50.55	1012	42.38	1018	
		$\mathcal{N}(0, 2^2)$	102.65	1012	84.89	1018	
TSP-500	Org	$\mathcal{N}(0, 4^2)$	184.40	1012	150.47	1018	
		$\mathcal{N}(0, 8^2)$	204.53	1010	171.14	1014	
		$\mathcal{U}(0,10)$	210.70	1012	175.41	1016	
		$\mathcal{N}(0,1)$	77.15	2826	60.56	2840	
		$\mathcal{N}(0, 2^2)$	157.12	2826	121.45	2839	
TSP-1000	Org	$\mathcal{N}(0,4^2)$	281.64	4224	214.55	2833	
		$\mathcal{N}(0,8^2)$	310.94	4218	243.77	4242	
		$\mathcal{U}(0,10)$	317.55	4218	249.28	4236	
		$\mathcal{N}(0,1)$	33.69	109	27.14	112	
		$\mathcal{N}(0,2^2)$	69.77	109	54.28	112	
	US	$\mathcal{N}(0,4^2)$	125.99	108	96.70	112	
		$\mathcal{N}(0,8^2)$	143.76	109	110.64	113	
		$\mathcal{U}(0,10)$	145.41	109	113.39	112	
		$\mathcal{N}(0,1)$	30.75	107	26.69	110	
		$\mathcal{N}(0,2^2)$	62.08	109	53.36	112	
TSP-200	CS	$\mathcal{N}(0,4^2)$	110.48	109	94.84	111	
		$\mathcal{N}(0,8^2)$	127.06	107	108.76	111	
		$\mathcal{U}(0,10)$	129.77	107	111.47	109	
		$\mathcal{N}(0,1)$	30.77	77	26.68	79	
		$\mathcal{N}(0, 2^2)$	61.87	76	53.38	79	
	CS-aligned	$\mathcal{N}(0, 4^2)$	110.99	78	94.59	79	
		$\mathcal{N}(0,8^2)$	126.69	76	108.42	79	
		$\mathcal{U}(0,10)$	129.28	76	111.49	78	
		$\mathcal{N}(0,1)$	59.81	1012	43.41	1020	
		$\mathcal{N}(0, 2^2)$	126.00	1013	86.76	1022	
	US	$\mathcal{N}(0, 4^2)$	237.72	1012	154.28	1022	
		$\mathcal{N}(0,8^2)$	261.79	1013	176.13	1022	
		$\mathcal{U}(0,10)$	263.66	1015	180.75	1022	

Table 6: Comparison of uniform sampling and our coreset method using TSP100-2D- $\mathcal{N}(0,1)$ as the training dataset on test data of varying sizes. We fix the sample size as 12951.

Continued on next page

TSP-500

TOD	Madad		Gree	edy	Greedy-	+2-opt
I SP size	SP size Method	lest distribution	Length (\downarrow)	Time (\downarrow)	Length (\downarrow)	Time (\downarrow)
		$\mathcal{N}(0,1)$	49.11	1012	42.25	1016
		$\mathcal{N}(0, 2^2)$	99.58	1012	84.60	1019
	CS	$\mathcal{N}(0, 4^2)$	178.56	1010	149.50	1016
		$\mathcal{N}(0,8^2)$	205.86	1012	170.54	1016
		$\mathcal{U}(0,10)$	208.36	1011	174.93	1016
		$\mathcal{N}(0,1)$	49.38	680	42.26	683
		$\mathcal{N}(0,2^2)$	99.66	680	84.52	684
	CS-aligned	$\mathcal{N}(0, 4^2)$	178.88	679	149.63	682
		$\mathcal{N}(0,8^2)$	204.42	678	170.62	682
		$\mathcal{U}(0,10)$	208.77	678	175.03	682
		$\mathcal{N}(0,1)$	94.71	2823	61.72	2848
		$\mathcal{N}(0,2^2)$	199.22	2825	123.56	2849
	US	$\mathcal{N}(0,4^2)$	382.77	4224	219.16	2847
		$\mathcal{N}(0,8^2)$	421.29	4218	249.08	4258
		$\mathcal{U}(0,10)$	426.61	4215	255.95	4254
		$\mathcal{N}(0,1)$	69.76	2823	59.59	2833
		$\mathcal{N}(0, 2^2)$	141.06	2825	119.64	2832
TSP-1000	CS	$\mathcal{N}(0, 4^2)$	252.92	4224	210.71	2832
		$\mathcal{N}(0,8^2)$	293.49	4218	240.42	4236
		$\mathcal{U}(0,10)$	299.80	4215	246.57	4234
		$\mathcal{N}(0,1)$	69.96	2825	59.57	2832
		$\mathcal{N}(0,2^2)$	140.95	2822	119.39	2833
	CS-aligned	$\mathcal{N}(0,4^2)$	253.63	2821	210.81	2830
		$\mathcal{N}(0,8^2)$	293.82	2815	240.11	2826
		$\mathcal{U}(0,10)$	300.03	2812	246.61	2827

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Table 7: Comparison of uniform sampling and our coreset method using TSP100-2D- $\mathcal{N}(0,1)$ as the training dataset on test data TSPLIB(Reinelt, 1991).

Commission	Mathad	Test distribution	Greedy		Greedy+2-opt	
Sample size	Method	Test distribution	Length (\downarrow)	Time (\downarrow)	Length (\downarrow)	Time (\downarrow)
128000	Org	$\mathcal{N}(0,1)$	129.35	108	112.23	106
	US	$\mathcal{N}(0,1)$	190.79	109	115.87	108
4003	CS	$\mathcal{N}(0,1)$	153.08	107	113.56	108
	CS-aligned	$\mathcal{N}(0,1)$	152.70	103	113.71	105
	US	$\mathcal{N}(0,1)$	166.40	106	114.47	108
8245	CS	$\mathcal{N}(0,1)$	140.49	107	113.04	106
	CS-aligned	$\mathcal{N}(0,1)$	140.18	104	112.91	104
	US	$\mathcal{N}(0,1)$	162.19	107	114.31	107
12951	CS	$\mathcal{N}(0,1)$	133.63	106	112.45	105
	CS-aligned	$\mathcal{N}(0,1)$	133.14	103	112.52	110

Results of TSP100-3D- $\mathcal{N}(0,1)$ Tables 8 to 10 show the results of TSP100-3D- $\mathcal{N}(0,1)$. Table 8 illustrates the overall acceleration improvement. For the test data without distribution shift, our method has comparable performance; for the test data occurring distribution shift, our method has better performance. Moreover, our coreset with alignment version performs better in the TSP-3D case. From Tables 9 and 10, alignment can perform better for higher dimension dataset (i.e., TSP-3D). Thus, the alignment version is promising for tackling high-dimensional data.

Method	Sample size	Labeling time	Coreset Time	Training Time	Total time
Org	128000	4940	-	30671	35611
	4103	160	-	2102	2262
US	7960	307	-	2729	3036
	12058	466	-	3514	3980
	4103	159	1012	2080	1379
CS	7960	309	1177	2712	4198
	12058	463	1675	3493	5631

Table 8: Time statistics for different phases of training on TSP100-3D- $\mathcal{N}(0, 1)$.

Table 10: Comparison of uniform sampling and our coreset method with training dataset TSP100-3D- $\mathcal{N}(0, 1)$ on test data of varying sizes. We fix the sample size as 12058.

TOD		The different disc	Gree	edy	A1'
I SP size	Method	lest distribution	Length (\downarrow)	Time (\downarrow)	Alignment time (\downarrow)
		$\mathcal{N}(0,1)$	30.02	77	-
		$\mathcal{N}(0, 2^2)$	61.03	76	-
TSP-200	Org	$\mathcal{N}(0, 4^2)$	109.78	77	-
		$\mathcal{N}(0, 8^2)$	126.15	76	-
		$\mathcal{U}(0,10)$	128.35	77	-
		$\mathcal{N}(0,1)$	48.66	682	-
		$\mathcal{N}(0,2^2)$	101.71	682	-
TSP-500	Org	$\mathcal{N}(0,4^2)$	184.94	682	-
		$\mathcal{N}(0,8^2)$	210.36	680	-
		$\mathcal{U}(0,10)$	212.62	683	-
		$\mathcal{N}(0,1)$	69.08	2828	-
		$\mathcal{N}(0, 2^2)$	144.10	2826	-
TSP-1000	Org	$\mathcal{N}(0,4^2)$	264.58	2824	-
		$\mathcal{N}(0,8^2)$	303.02	2821	-
		$\mathcal{U}(0,10)$	313.23	2818	-
		$\mathcal{N}(0,1)$	32.30	76	-
		$\mathcal{N}(0,2^2)$	67.81	77	-
	US	$\mathcal{N}(0,4^2)$	130.76	77	-
		$\mathcal{N}(0,8^2)$	152.74	77	-
		$\mathcal{U}(0,10)$	153.90	77	-
		$\mathcal{N}(0,1)$	32.72	76	-
		$\mathcal{N}(0,2^2)$	66.08	76	-
TSP-200	CS	$\mathcal{N}(0,4^2)$	120.64	78	-
		$\mathcal{N}(0,8^2)$	139.91	77	-
		$\mathcal{U}(0,10)$	142.84	77	-
		$\mathcal{N}(0,1)$	33.25	76	2
		$\mathcal{N}(0, 2^2)$	69.79	75	5
	CS-aligned	$\mathcal{N}(0, 4^2)$	118.44	77	7
		$\mathcal{N}(0,8^2)$	135.90	77	10
		$\mathcal{U}(0,10)$	138.27	76	12
		$\mathcal{N}(0,1)$	58.45	682	
		$\mathcal{N}(0, 2^2)$	127.53	681	-
	US	$\mathcal{N}(0, 4^2)$	264.93	682	-
		$\mathcal{N}(0,8^2)$	316.73	681	-
		$\mathcal{U}(0,10)$	318.39	680	
		$\mathcal{N}(0,1)$	59.69	682	-

Continued on next page TSP-500 CS

CS

		I I I I I I I I I I I I I I I I I I I	Gree	dv	
TSP size	Method	Test distribution	Length (\downarrow)	Time (\downarrow)	
		$\mathcal{N}(0, 2^2)$	122.24	681	-
		$\mathcal{N}(0, 4^2)$	231.96	682	-
		$\mathcal{N}(0, 8^2)$	266.08	677	-
		$\mathcal{U}(0,10)$	268.75	680	-
		$\mathcal{N}(0,1)$	56.47	682	16
		$\mathcal{N}(0,2^2)$	121.33	681	19
	CS-aligned	$\mathcal{N}(0, 4^2)$	205.45	682	23
		$\mathcal{N}(0,8^2)$	238.14	681	26
		$\mathcal{U}(0,10)$	245.01	681	30
	US	$\mathcal{N}(0,1)$	86.01	2828	-
		$\mathcal{N}(0,2^2)$	191.39	2828	-
		$\mathcal{N}(0, 4^2)$	397.40	2825	-
		$\mathcal{N}(0,8^2)$	470.55	2820	-
		$\mathcal{U}(0,10)$	550.11	2819	-
		$\mathcal{N}(0,1)$	86.57	2826	-
		$\mathcal{N}(0,2^2)$	181.84	2827	-
TSP-1000	CS	$\mathcal{N}(0, 4^2)$	342.64	2822	-
		$\mathcal{N}(0,8^2)$	395.66	2820	-
		$\mathcal{U}(0,10)$	441.82	2818	-
		$\mathcal{N}(0,1)$	86.87	2828	236
		$\mathcal{N}(0,2^2)$	186.88	2823	433
	CS-aligned	$\mathcal{N}(0,4^2)$	320.68	2825	624
		$\mathcal{N}(0,8^2)$	372.34	2820	861
		$\mathcal{U}(0,10)$	383.92	2819	980

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Then, we take TSP-3D as an example to compare the performance of the random heuristic alignment method (CS-rand-aligned) with our proposed alignment method (CS-aligned).

As shown in Table 11, the results demonstrate that the CS-rand-aligned method performs similarly to the unaligned approach, providing little improvement. In contrast, our alignment method significantly enhances performance, confirming its practical effectiveness.

Table 11: Comparison of rand alignment (CS-rand-aligned) and our alignment (CS-aligned) method with training dataset TSP100-3D- $\mathcal{N}(0,1)$ on test data of varying sizes. We fix the sample size as 12058.

TSP size	Method	Test distribution	Greedy Length (\downarrow) Time (\downarrow)		Alignment time (\downarrow)
	US	$\mathcal{U}(0,10)$	100.41	37	-
TSD 100	CS	$\mathcal{U}(0,10)$	95.27	36	-
131-100	CS-aligned	$\mathcal{U}(0,10)$	94.13	37	3
	CS-rand-aligned	$\mathcal{U}(0,10)$	95.62	37	11
	US	$\mathcal{U}(0,10)$	153.90	77	-
TSD 200	CS	$\mathcal{U}(0,10)$	142.84	77	-
131-200	CS-aligned	$\mathcal{U}(0,10)$	138.27	76	12
	CS-rand-aligned	$\mathcal{U}(0,10)$	144.84	75	17
	US	$\mathcal{U}(0,10)$	318.39	680	-
TSD 500	CS	$\mathcal{U}(0,10)$	268.75	680	-
131-300	CS-aligned	$\mathcal{U}(0,10)$	245.01	681	30
	CS-rand-aligned	$\mathcal{U}(0,10)$	255.92	674	23
	US	$\mathcal{U}(0,10)$	550.11	2819	-
TSP-1000	CS	$\mathcal{U}(0,10)$	441.82	2826	-
151-1000	CS-aligned	$\mathcal{U}(0,10)$	383.92	2818	980

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	– continued fr	om previous page			
TSP size	Method	Test distribution	Gree Length (\downarrow)	edy Time (\downarrow)	
	CS-rand-aligned	$\mathcal{U}(0,10)$	429.90	2817	493

Results of TSP100-2D- $\mathcal{U}(0, 10)$ Tables 12 to 15 present the results on training dataset TSP100-2D- $\mathcal{U}(0, 10)$. The training datasets are generated by the uniform sampling and our coreset technique respectively. Table 12 clearly demonstrates the overall acceleration improvement. All the results show that both our method and uniform sampling method perform better as the sample size increases. For both the test data with distribution shift and without distribution shift, our method consistently shows better performance. Table 14 further illustrates that our method can generalize to large-scale TSP problems better. Table 15 confirms that our method outperforms other approaches on the TSPLIB dataset.

Results of TSP100-3D- $\mathcal{U}(0, 10)$ Tables 16 and 17 present the results on training dataset TSP100-2D- $\mathcal{U}(0, 10)$.

E FULL EXPERIMENTS WITH MIS

Dataset For Maximal Independent Set (MIS), we train our model on ER-[90-100] dataset (Erd6s & Rényi, 1960), where ER-[n-N] means that the graph contains n to N nodes. We set the connection probability as 0.15 as in (Sun & Yang, 2023). The labels of our MIS datasets are obtained by using the KaMIS2 solver. Our training dataset of MIS consists of 128000 ER-[90-100] instances. We evaluate our method on ER-[90-100], ER-[400-500], ER-[900-1000] and SATLIB.

Setting We use the DIFUSCO (Sun & Yang, 2023) as our NCO solver. To quantify the performance of different methods, we use two criteria: the average size of the independent set (Size) and the total runtime (Time). The term "Greedy" refers to the greedy decoding method of DIFUSCO, and "Sampling" is the sampling decoding. The terms US and CS represent uniform sampling and our coreset method, respectively. We use the results on the full dataset with 128000 data as a baseline.

Results of MIS Tables 18, 20 and 27 present the results on training dataset ER-[90-100]. The training datasets are generated by the uniform sampling and our coreset technique respectively. Table 18 clearly demonstrates the overall acceleration improvement. All the results show that both our method and uniform sampling method perform better as the sample size increases. Table 27 demonstrates that our method can generalize to large-scale ER problems better. Table 20 show that the performance of our method and uniform sampling method is similar on SATLIB.

F FULL EXPERIMENTS WITH CVRP

In this section, we take CVRP100 as an example to show the advantages of our coreset method. All experiments of CVRP are conducted on NVIDIA GeForce RTX 3090 GPU. We take the NCO (Luo et al., 2023) as our backbone solver, and set its training epoch as 20.

Dataset We apply our method on CVRP100 Euclidean instances. The labels of CVRP100 instances are obtained by using the LKH-3 heuristic solver (Helsgaun, 2017); each coordinate of the nodes in a CVRP instance is generated by x%1, where x is randomly sampled either from a normal distribution $\mathcal{N}(0, \sigma^2)$ or uniform sampling $\mathcal{U}(0, 1)$. Our training data CVRP100- $\mathcal{N}(0, 0.1^2)$ consists of 125,000 instances generated by the normal distribution $\mathcal{N}(0, 0.1^2)$ and 3000 instances by the uniform distribution $\mathcal{U}(0, 1)$. While the training dataset CVRP100- $\mathcal{U}(0, 1)$ consists of 128000 instances generated by uniform distribution $\mathcal{U}(0, 1)$.

Indeed, the distributions of the training dataset CVRP100- $\mathcal{N}(0, 0.1^2)$ is very close to the normal distribution $\mathcal{N}(0, 0.1^2)$. Hence, we regard the test data sampled from distribution $\mathcal{N}(0, 0.1^2)$ as having no distribution shift. The test data are sampled from a single distribution, either a normal distribution or uniform distribution. Specifically, we sample 1280 test data for CVRP100, and 128

$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Sample size Method		Test distribution	Gree Length (\downarrow)	edy Time (\downarrow)	Alignment time (\downarrow)
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{N}(0,1)$	20.80	364	-
$\begin{array}{c c c c c c c c c c c c c c c c c c c $			$\mathcal{N}(0, 2^2)$	42.25	366	-
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	128000	Org	$\mathcal{N}(0, 4^2)$	76.57	362	-
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $			$\mathcal{N}(0, 8^2)$	88.45	360	-
$\begin{array}{c c c c c c c c c c c c c c c c c c c $			$\mathcal{U}(0, 10)$	90.55	358	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{N}(0,1)$	24.92	480	-
$\begin{array}{c c c c c c c c c c c c c c c c c c c $			$\mathcal{N}(0,2^2)$	49.96	482	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$		US	$\mathcal{N}(0,4^2)$	96.60	483	-
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $			$\mathcal{N}(0,8^2)$	116.65	482	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{U}(0,10)$	119.78	481	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{N}(0,1)$	24.89	364	-
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	4102	~~	$\mathcal{N}(0, 2^2)$	50.46	384	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	4103	CS	$\mathcal{N}(0, 4^2)$	106.35	360	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{N}(0, 8^2)$	109.12	360	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{U}(0,10)$	111.63	353	-
$\begin{array}{c cccc} & & & & & & & & & & & & & & & & & $			$\mathcal{N}(0,1)$	23.36	479	2
$\begin{tabular}{ c c c c c c c c c c c c c c c c c c c$			$\mathcal{N}(0,2^2)$	47.00	480	4
$\begin{array}{c c c c c c c c c c c c c c c c c c c $		CS-aligned	$\mathcal{N}(0, 4^2)$	91.94	479	7
$\frac{\mathcal{U}(0,10)}{\mathcal{N}(0,1)} = \frac{108.81}{23.62} = \frac{483}{477} = \frac{11}{\sqrt{\mathcal{N}(0,2^2)}} = \frac{48.32}{48.32} = \frac{477}{477} = \frac{11}{\sqrt{\mathcal{N}(0,2^2)}} = \frac{48.32}{48.32} = \frac{477}{477} = \frac{11}{\sqrt{\mathcal{N}(0,2^2)}} = \frac{48.32}{477} = \frac{11}{\sqrt{\mathcal{N}(0,2^2)}} = \frac{48.4}{477} = \frac{11}{\sqrt{\mathcal{N}(0,10)}} = \frac{115.62}{481} = \frac{481}{7} = \frac{11}{\sqrt{\mathcal{N}(0,10)}} = \frac{115.62}{481} = \frac{481}{7} = \frac{11}{\sqrt{\mathcal{N}(0,10)}} = \frac{115.62}{481} = \frac{11}{7} = \frac{11}{\sqrt{\mathcal{N}(0,12^2)}} = \frac{47.10}{47.10} = \frac{361}{61} = \frac{11}{7} = \frac{11}{\sqrt{\mathcal{N}(0,12^2)}} = \frac{47.10}{47.10} = \frac{11}{361} = \frac{11}{7} = \frac{11}{\sqrt{\mathcal{N}(0,12^2)}} = \frac{11}{47} = \frac{11}{\sqrt{\mathcal{N}(0,12^2)}} = \frac{11}{483} = \frac{11}{10} = \frac{11}{\sqrt{\mathcal{N}(0,12^2)}} = \frac{11}{48.10} = \frac{11}{10} = \frac{11}{\sqrt{\mathcal{N}(0,12^2)}} = \frac{11}{48.10} = \frac{11}{10} = \frac{11}{\sqrt{\mathcal{N}(0,12^2)}} = \frac{11}{48.10} = \frac{11}{10} = \frac{11}{10} = \frac{11}{100.42} = \frac{11}{100.4$			$\mathcal{N}(0,8^2)$	106.50	482	9
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{U}(0,10)$	108.81	483	11
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{N}(0,1)$	23.62	477	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		TIC.	$\mathcal{N}(0, 2^2)$	48.32	477	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		US	$\mathcal{N}(0, 4^2)$	92.92	484	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{N}(0, 8^2)$	111.98	479	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{U}(0,10)$	115.62	481	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{N}(0,1)$	23.41	362	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	7060	<u> </u>	$\mathcal{N}(0,2^2)$	47.10	361	-
$12058 \begin{array}{ c c c c c c c c c c c c c c c c c c c$	7900	CS	$\mathcal{N}(0, 4^2)$	86.20	362	-
$12058 \begin{array}{ c c c c c c c c c c c c c c c c c c c$			$\mathcal{N}(0, 8^2)$	99.50	365	-
$12058 \begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{U}(0, 10)$	101.25	359	-
$12058 \begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{N}(0,1)$	22.83	476	12
$12058 \begin{array}{ c c c c c c c c c c c c c c c c c c c$		CC alianad	$\mathcal{N}(0, 2^2)$	46.06	4/4	14
$12058 \begin{array}{c ccccccccccccccccccccccccccccccccccc$		CS-aligned	$\mathcal{N}(0, 4^2)$	85.71	483	16
$12058 \begin{array}{ c c c c c c c c c c c c c c c c c c c$			$\mathcal{N}(0, 8^2)$	97.01	480	1/
$12058 \begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\frac{\mathcal{U}(0,10)}{\mathcal{N}(0,1)}$	99.10	481	19
$12058 \qquad \begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{N}(0, 1)$ $\mathcal{N}(0, 2^2)$	45.50	300	-
$12058 \qquad \begin{array}{c ccccccccccccccccccccccccccccccccccc$		US	$\mathcal{N}(0, 2)$ $\mathcal{N}(0, 4^2)$	43.32	268	-
$12058 \qquad \begin{array}{c ccccccccccccccccccccccccccccccccccc$		05	$\mathcal{N}(0,4)$ $\mathcal{N}(0,8^2)$	04.20	308	-
$12058 \qquad \begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{N}(0, \delta)$	98.37	372	-
$12058 \qquad CS \qquad \begin{array}{c} \mathcal{N}(0,1) & 22.10 & 371 & \mathbf{-} \\ \mathcal{N}(0,2^2) & 44.40 & 362 & \mathbf{-} \\ \mathcal{N}(0,4^2) & 80.47 & 361 & \mathbf{-} \\ \mathcal{N}(0,8^2) & 93.00 & 361 & \mathbf{-} \\ \mathcal{U}(0,10) & 95.25 & 362 & \mathbf{-} \\ \hline \mathcal{N}(0,1) & 22.30 & 396 & 21 \\ \mathcal{N}(0,2^2) & 44.80 & 371 & 22 \\ CS\text{-aligned} & \mathcal{N}(0,4^2) & 79.96 & 388 & 23 \\ \mathcal{N}(0,8^2) & 92.28 & 377 & 25 \\ \mathcal{U}(0,10) & 94.13 & 366 & 26 \end{array}$			$\frac{\mathcal{U}(0,10)}{\mathcal{N}(0,1)}$	22 10	307	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\mathcal{N}(0, 1)$ $\mathcal{N}(0, 2^2)$	22.10 11 10	371	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	12058	CS	$\mathcal{N}(0, 2)$	44.4V 80 47	361	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	1_000	0	N(0, 4)	02.00	261	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$			$\frac{1}{10}(0, 0)$	95.00	367	-
$\begin{array}{cccccccccccccccccccccccccccccccccccc$			$\frac{\mathcal{U}(0, 10)}{\mathcal{N}(0, 1)}$	22.25	302	- 21
CS-aligned $\mathcal{N}(0, 2^2)$ 79.96 388 23 $\mathcal{N}(0, 8^2)$ 92.28 377 25 $\mathcal{U}(0, 10)$ 94.13 366 26			$\mathcal{N}(0, 1)$ $\mathcal{N}(0, 2^2)$	22.30 44 80	371	$\frac{21}{22}$
$\begin{array}{c} \mathcal{N}(0,4^{2}) & 77.70 & 566 & 25 \\ \mathcal{N}(0,8^{2}) & 92.28 & 377 & 25 \\ \mathcal{U}(0,10) & \mathbf{9413} & 366 & 26 \end{array}$		CS-aligned	$\mathcal{N}(0, 2)$	70 0 4	399	22
$\mathcal{U}(0,0)$ 72.20 $\mathcal{I}(1)$ 23 $\mathcal{U}(0,10)$ 94 13 366 26		C5 anglieu	$\mathcal{N}(0, 4)$	17.70	300 277	23 25
· · · · · · · · · · · · · · · · · · ·			$\frac{1}{1}(0, 0, 0)$	94 13	366	25 26

Table 9: Comparison of uniform sampling and our coreset method using TSP100-3D- $\mathcal{N}(0,1)$ as the training dataset on test data TSP100-3D from different distributions.

Method	Sample size	Labeling time	Coreset Time	Training Time	Total time
Org	128000	4883	-	27686	32569
	4003	153	-	2387	2540
US	8245	315	-	3217	3532
	12951	495	-	3578	4073
	4003	154	859	2331	3344
CS	8245	315	841	3217	4373
	12951	493	1293	3516	5302

Table 12: Time statistics for different phases of training on TSP100-2D- $\mathcal{U}(0, 10)$.

Table 13: Comparison of uniform sampling and our coreset method using TSP100-2D- $\mathcal{U}(0, 10)$ as the training dataset on test data TSP100-2D from different distributions.

Sample size	Mathod	Test distribution	Gree	edy	Greedy+2-opt	
Sample size	Method	Test distribution	Length (\downarrow)	Time (\downarrow)	Length (\downarrow)	Time (\downarrow)
128000	Org	$\mathcal{U}(0,10)$	86.77	358	79.53	353
	US	$\mathcal{U}(0,10)$	100.62	355	80.97	375
4003	CS	$\mathcal{U}(0,10)$	97.69	461	80.94	383
	CS-aligned	$\mathcal{U}(0,10)$	97.71	363	80.95	369
	US	$\mathcal{U}(0,10)$	92.66	352	80.33	365
8245	CS	$\mathcal{U}(0,10)$	93.12	381	80.38	372
	CS-aligned	$\mathcal{U}(0,10)$	93.27	351	80.41	354
	US	$\mathcal{U}(0,10)$	91.98	362	80.34	367
12951	CS	$\mathcal{U}(0,10)$	92.33	354	80.23	356
	CS-aligned	$\mathcal{U}(0,10)$	92.39	360	80.28	367

Table 14: Comparison of uniform sampling and our coreset method using TSP100-2D- $\mathcal{U}(0, 10)$ as the training dataset on test data of varying sizes. We fix the sample size as 12951.

TCD size	Mathad	Test distribution	Gree	edy	Greedy+2-opt	
I SP Size	Method	Test distribution	Length (\downarrow)	Time (\downarrow)	Length (\downarrow)	Time (\downarrow)
TSP-200	Org	$\mathcal{U}(0,10)$	125.06	72	110.89	73
TSP-500	Org	$\mathcal{U}(0,10)$	195.68	675	172.19	677
TSP-1000	Org	$\mathcal{U}(0,10)$	276.69	2815	241.59	2824
	US	$\mathcal{U}(0,10)$	130.05	76	111.71	81
TSP-200	CS	$\mathcal{U}(0,10)$	129.52	75	111.31	81
	CS-aligned	$\mathcal{U}(0,10)$	129.95	77	111.35	76
	US	$\mathcal{U}(0,10)$	238.46	677	177.30	682
TSP-500	CS	$\mathcal{U}(0,10)$	221.76	677	175.94	682
	CS-aligned	$\mathcal{U}(0,10)$	221.72	675	175.89	683
	US	$\mathcal{U}(0,10)$	434.16	2811	254.37	2837
TSP-1000	CS	$\mathcal{U}(0,10)$	346.71	2811	251.40	2831
	CS-aligned	$\mathcal{U}(0,10)$	347.46	2813	251.35	2833

Samula siza	Mathad	Test distribution	Gree	edy	Greedy-	Greedy+2-opt	
Sample size	Method	Test distribution	Length (\downarrow)	Time (\downarrow)	Length (\downarrow)	Time (\downarrow)	
128000	Org	$\mathcal{N}(0,1)$	125.39	102	110.90	104	
	US	$\mathcal{U}(0,10)$	183.71	104	115.21	110	
4003	CS	$\mathcal{U}(0,10)$	164.81	106	114.50	108	
	CS-aligned	$\mathcal{U}(0,10)$	166.04	105	114.41	103	
	US	$\mathcal{U}(0,10)$	161.75	107	113.69	107	
8245	CS	$\mathcal{U}(0,10)$	132.06	103	111.52	111	
	CS-aligned	$\mathcal{U}(0,10)$	145.90	103	112.86	103	
	US	$\mathcal{U}(0,10)$	157.10	104	113.98	105	
12951	CS	$\mathcal{U}(0,10)$	140.97	104	112.96	106	
	CS-aligned	$\mathcal{U}(0,10)$	141.81	501	112.81	110	

Table 15: Comparison of uniform sampling and our coreset method using TSP100-2D- $\mathcal{U}(0, 10)$ as the training dataset on test data TSPLIB(Reinelt, 1991).

Table 16: Comparison of uniform sampling and our coreset method using TSP100-3D- $\mathcal{U}(0, 10)$ as the training dataset on test data TSP100-3D from different distributions.

Sample size	Mathad	Test distribution	Gree	Greedy		
Sample size	Method	Test distribution	Length (\downarrow)	Time (\downarrow)		
128000	Org	$\mathcal{U}(0,10)$	87.84	350		
	US	$\mathcal{U}(0,10)$	108.74	350		
4122	CS	$\mathcal{U}(0,10)$	108.45	352		
	CS-aligned	$\mathcal{U}(0,10)$	106.11	381		
	US	$\mathcal{U}(0,10)$	95.78	360		
8245	CS	$\mathcal{U}(0,10)$	98.53	347		
	CS-aligned	$\mathcal{U}(0,10)$	97.89	358		
	US	$\mathcal{U}(0,10)$	93.38	364		
12951	CS	$\mathcal{U}(0,10)$	93.15	348		
	CS-aligned	$\mathcal{U}(0,10)$	92.91	354		

Table 17: Comparison of uniform sampling and our coreset method using TSP100-3D- $\mathcal{U}(0, 10)$ as the training dataset on test data of varying sizes. We fix the sample size as 12951.

TSD size	Mathad	Test distribution	Greedy		
I SP Size	Method	Test distribution	Length (\downarrow)	Time (\downarrow)	
TSP-200	Org	$\mathcal{U}(0,10)$	125.98	73	
TSP-500	Org	$\mathcal{U}(0,10)$	210.15	674	
TSP-1000	Org	$\mathcal{U}(0,10)$	313.00	2818	
	US	$\mathcal{U}(0,10)$	145.67	78	
TSP-200	CS	$\mathcal{U}(0,10)$	144.27	76	
	CS-aligned	$\mathcal{U}(0,10)$	143.46	75	
	US	$\mathcal{U}(0,10)$	302.16	674	
TSP-500	CS	$\mathcal{U}(0,10)$	244.63	676	
	CS-aligned	$\mathcal{U}(0,10)$	236.33	689	
-	US	$\mathcal{U}(0,10)$	535.90	2815	
TSP-1000	CS	$\mathcal{U}(0,10)$	367.04	2812	
	CS-aligned	$\mathcal{U}(0,10)$	308.71	2816	

	Table 1	o. Thic statistics	s for unreferit ph	ases of MID.	
Method	Sample size	Labeling time	Coreset Time	Training Time	Total time
Org	128000	102600	-	16762	119362
	3973	3184	-	1726	4910
US	8001	6413	-	2503	8916
	12417	9953	-	3400	13353
	3973	4007	2263	3227	9497
CS	8001	7322	2424	5031	14777
	12417	11032	2511	7011	20554

Table 18: Time statistics for different phases of MIS.

Table 19: Comparison of uniform sampling and our coreset method with training dataset ER-[90-100] on test data from different distributions.

Sample size	Mathad	Test distribution	Gre	Greedy		Sampling	
Sample size	Method	Test distribution	Size (\uparrow)	Time (\downarrow)	Size (\uparrow)	Time (\downarrow)	
		ER-[v90-100]	22.34	64	23.27	70	
128000	Org	ER-[400-500]	34.66	134	36.69	551	
		ER-[700-800]	37.51	393	40.06	1463	
		ER-[v90-100]	19.57	64	21.34	319	
	US	ER-[400-500]	27.84	134	29.48	552	
3073		ER-[700-800]	30.30	391	32.23	1469	
3913		ER-[v90-100]	19.77	63	21.49	72	
	CS	ER-[400-500]	27.99	135	29.85	558	
		ER-[700-800]	30.61	392	32.38	1524	
		ER-[v90-100]	20.38	64	22.33	70	
	US	ER-[400-500]	28.81	135	30.59	554	
8001		ER-[700-800]	31.28	393	32.77	1471	
8001	-	ER-[v90-100]	20.25	64	22.27	73	
	CS	ER-[400-500]	29.59	134	31.91	549	
		ER-[700-800]	32.23	393	33.93	1460	
		ER-[v90-100]	20.80	63	22.30	71	
	US	ER-[400-500]	30.36	135	32.55	549	
12417		ER-[700-800]	32.30	393	34.25	1458	
1241/		ER-[v90-100]	21.04	63	22.55	70	
	CS	ER-[400-500]	31.00	135	33.11	548	
		ER-[700-800]	32.96	393	35.59	1462	

Table 20: Comparison of uniform sampling and our coreset method with training dataset ER-[90-100] on test data SATLIB.

Samala siza	Mathad	athad Tast distribution		Greedy		Sampling	
Sample size	Method	Test distribution	Size (\uparrow)	Time (\downarrow)	Size (\uparrow)	Time (\downarrow)	
128000	Org	ER-[v90-100]	22.34	64	23.27	70	
2072	US	SATLIB	1 015.64	120	1022.81	490	
3913	CS	SATLIB	1015.31	122	1021.24	518	
0245	US	SATLIB	410.61	296	413.49	898	
8245	CS	SATLIB	409.87	299	413.34	902	
10/17	US	SATLIB	410.32	304	413.55	906	
12417	CS	SATLIB	410.83	297	414.04	900	

Sample size	Method	Test distribution	RRC-budget (\downarrow)	Length (\downarrow)	$\text{Gap}\;(\downarrow)$	Time (\downarrow)
			0	17.64	6.34%	3
			50	16.58	-0.02%	45
128000	Org	CVRP100	100	16.43	-0.96%	86
			200	16.30	-1.70%	162
			500	16.18	-2.43%	397
			0	19.56	17.91%	3
			50	17.85	7.64%	45
	US	CVRP100	100	17.57	5.96%	86
			200	17.30	4.28%	162
1137			500	17.03	2.71%	397
4437			0	19.16	15.51%	3
		CVRP100	50	17.65	6.41%	45
	CS		100	17.38	4.78%	86
			200	17.15	3.38%	163
			500	16.91	2.01%	400
		CVRP100	0	18.77	13.18%	3
	US		50	17.35	4.62%	47
			100	17.13	3.27%	91
			200	16.92	2.01%	172
8082			500	16.71	0.75%	423
0002			0	18.49	11.51%	3
		CVRP100	50	17.19	3.66%	45
	CS		100	16.98	2.38%	88
			200	16.78	1.19%	168
			500	16.60	0.06%	413
			0	18.65	12.47%	3
			50	17.24	3.95%	47
	US	CVRP100	100	17.01	2.59%	90
			200	16.82	1.43%	170
12175			500	16.63	0.25%	415
12115			0	18.53	11.72%	3
			50	17.18	3.56%	47
	CS	CVRP100	100	16.96	2.23%	90
			200	16.77	1.13%	170
			500	16.58	-0.01%	415

Table 21: Comparison of uniform sampling and our coreset method with training dataset CVRP100- $\mathcal{N}(0,0.1^2)$ on test data from different distributions.

test data for other cases. Obviously, the uniform distribution has the highest entropy and thus the highest diversity. For these Gaussian distributions, the larger the variance, the larger the diversity.

Results of CVRP100 Tables 21 to 23 present the results on training dataset CVRP100- $\mathcal{N}(0, 0.1^2)$. The training datasets are generated by the uniform sampling and our coreset technique respectively. The results show that both our method and uniform sampling method perform better as the sample size increases. For both the test data with distribution shift and without distribution shift, our method consistently shows better performance. Table 22 further illustrates that our method can generalize to large-scale CVRP problems better. Table 23 shows the performance on the CVRPLIB dataset.

Tables 24 to 26 present the results on training dataset CVRP100- $\mathcal{U}(0, 1)$. The training datasets are generated by the uniform sampling and our coreset technique respectively. The results show that both our method and uniform sampling method perform better as the sample size increases. For both the test data with distribution shift and without distribution shift, our method consistently shows better performance. Table 25 further illustrates that our method can generalize to large-scale CVRP problems better. Table 26 shows the performance on the CVRPLIB dataset.

Table 22: Comparison of uniform sampling and our coreset method with training dataset CVRP100- $\mathcal{N}(0, 0.1^2)$ on test data of varying sizes. We fix the sample size as 12175.

Sample size	Method	Test distribution	RRC-budget (\downarrow)	Length (\downarrow)	$\text{Gap}\;(\downarrow)$	Time (\downarrow)
			0	22.70	12.52%	2
			50	21.50	6.60%	26
		CVRP200	100	21.34	5.77%	49
			200	21.16	4.90%	105
			500	20.96	3.88%	256
			0	41.68	11.95%	2
			50	40.23	8.06%	179
128000	Org	CVRP500	100	39.91	7.20%	298
			200	39.64	6.48%	604
			500	39.32	5.62%	1628
			0	44.62	20.29%	73
			50	42.74	15.25%	895
		CVRP1000	100	42.36	14.22%	1810
			200	41.81	12.72%	3672
			500	41.19	11.04%	9108
			0	24.17	19.80 %	2
		CVRP200	50	22.79	12.97%	26
	US		100	22.52	11.61%	50
			200	22.24	10.27%	108
10175			500	21.89	8.52%	263
12175			0	24.00	18.97%	2
	CS		50	22.60	12.03%	25
		CVRP200	100	22.36	10.83%	48
			200	22.13	9.71%	105
			500	21.75	7.84%	256
			0	45.23	21.50%	2
			50	43.40	16.58%	173
	US	CVRP500	100	42.84	15.06%	288
12175			200	42.39	13.85%	584
			500	41.82	12.32%	1579
12175			0	45.10	21.15%	2
			50	43.19	16.00%	173
	CS	CVRP500	100	42.73	14.78%	287
			200	42.28	13.56%	583
			500	41.64	11.85%	1576

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Sample size	Method	Test distribution	RRC-budget (\downarrow)	Length (\downarrow)	$\operatorname{Gap}(\downarrow)$	Time (\downarrow)
-			0	53.01	42.91%	74
			50	49.42	33.23%	897
	US	CVRP1000	100	48.81	31.60%	1819
			200	47.95	29.28%	3708
12175			500	46.91	26.48%	9180
12175			0	51.88	39.88%	73
	CS	CVRP1000	50	48.24	30.06%	892
			100	47.71	28.64%	1805
			200	47.00	26.71%	3672
			500	46.12	24.34%	9108
			200 500	47.00 46.12	26.71% 24.34%	3672 9108

Table 22: Comparison of uniform sampling and our coreset method with training dataset CVRP100- $\mathcal{N}(0,0.1^2)$ on test data of varying sizes. We fix the sample size as 12175.

Table 23: Comparison of uniform sampling and our coreset method with training dataset CVRP100- $\mathcal{N}(0,0.1^2)$ on test data CVRPLib.

Sample size	Method	Test distribution	RRC-budget (\downarrow)	Length (\downarrow)	$\text{Gap}\;(\downarrow)$	Time (\downarrow)
			0	913.17	16.48%	1
			50	787.20	0.41%	4
128000	Org	CVRPLib	100	787.20	0.41%	7
	U		200	787.20	0.41%	12
			500	787.20	0.41%	29
			0	927.95	18.36%	1
			50	868.66	10.80%	4
	US	CVRPLib	100	864.51	10.27%	7
			200	844.59	7.73%	14
1127			500	844.59	7.73%	33
4437			0	950.06	21.18%	1
			50	841.93	7.39%	4
	CS	CVRPLib	100	837.50	6.82%	7
			200	837.50	6.82%	14
			500	834.28	6.41%	32
			0	971.82	23.96%	1
		CVRPLib	50	867.72	10.68%	4
	US		100	796.05	1.54%	7
			200	796.05	1.54%	13
0007			500	787.81	0.49%	30
8082			0	917.07	16.97%	1
			50	829.60	5.82%	4
	CS	CVRPLib	100	829.60	5.82%	7
			200	823.85	5.08%	13
			500	791.81	1.00%	30
			0	1016.27	29.63 %	1
			50	865.31	10.37%	4
	US	CVRPLib	100	865.31	10.37%	7
10175			200	858.44	9.49%	13
			500	858.44	9.49%	30
12175			0	917.09	16.98%	1
			50	811.18	3.47%	4
	CS	CVRPLib	100	810.27	3.35%	7
			200	806.12	2.82%	13
			500	806.12	2.82%	30

Sample size	Method	Test distribution	RRC-budget (\downarrow)	Length (\downarrow)	$\text{Gap}\;(\downarrow)$	Time (\downarrow)
			0	17.35	4.64%	3
			50	16.36	-1.38%	47
128000	Org	CVRP100	100	16.22	-2.18%	90
			200	16.13	-2.75%	170
			500	16.02	-3.40%	418
			0	18.48	11.43 %	3
			50	17.15	3.39%	44
	US	CVRP100	100	16.93	2.08%	85
			200	16.74	0.93%	161
4607			500	16.57	-0.10%	396
4097			0	18.44	11.20%	3
			50	17.10	3.10%	44
	CS	CVRP100	100	16.89	1.85%	85
			200	16.70	0.69%	161
			500	16.53	-0.34%	
			0	18.22	9.83%	3
		CVRP100	50	16.91	1.96%	44
	US		100	16.71	0.78%	83
			200	16.55	-0.21%	158
7694			500	16.39	-1.17%	389
7094			0	18.16	9.52%	3
		CVRP100	50	16.91	1.97%	45
	CS		100	16.72	0.82%	86
			200	16.56	-0.18%	162
			500	16.40	-1.10%	400
			0	18.03	8.70%	3
			50	16.82	1.41%	45
	US	CVRP100	100	16.64	0.34%	86
12033			200	16.50	-0.54%	162
			500	16.35	-1.43%	395
12055			0	18.01	8.59%	3
			50	16.81	1.36%	44
	CS	CVRP100	100	16.62	0.21%	84
			200	16.48	-0.64%	160
			500	16.34	-1.55%	39

Table 24: Comparison of uniform sampling and our coreset method with training dataset CVRP100- $\mathcal{U}(0,1)$ on test data from different distributions.

Table 25: Comparison of uniform sampling and our coreset method with training dataset CVRP100- $\mathcal{U}(0,1)$ on test data of varying sizes. We fix the sample size as 12033.

Sample size	Method	Test distribution	RRC-budget (\downarrow)	Length (\downarrow)	$\text{Gap}\;(\downarrow)$	Time (\downarrow)
			0	22.41	11.08%	2
			50	21.20	5.08%	36
		CVRP200	100	21.05	4.36%	68
			200	20.93	3.76%	145
			500	20.75	2.88%	353
			0	41.01	10.16%	1
			50	39.69	6.61%	190
128000	Org	CVRP500	100	39.42	5.87%	319
			200	39.16	5.20%	645
			500	38.91	4.51%	1730
			0	43.09	16.19%	73

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CVRP1000

Sample size	Method	Test distribution	RRC-budget (\downarrow)	Length (\downarrow)	$\operatorname{Gap}(\downarrow)$	Time (\downarrow)
			50	37.09	11.85%	904
			100	41.17	11.00%	1828
			200	40.76	9.90%	3708
			500	40.29	8.63%	9216
			0	23.29	15.45%	2
			50	21.98	8.94%	28
	US	CVRP200	100	21.75	7.82%	54
			200	21.57	6.91%	115
12033			500	21.33	5.72%	281
12033			0	20.17	14.97%	2
			50	21.93	8.70%	29
	CS	CVRP200	100	21.75	7.83%	54
			200	21.59	7.04%	116
			500	21.30	5.50%	282
			0	37.23	16.76%	2
			50	41.77	12.19%	177
	US	CVRP500	100	41.38	11.15%	296
			200	41.02	10.18%	598
12022			500	40.51	8.82%	1620
12055			0	43.19	16.02%	2
	CS		50	41.66	11.89%	176
		CVRP500	100	41.25	10.81%	295
			200	40.88	9.80%	597
			500	40.40	8.52%	1609
			0	48.20	29.96%	73
			50	46.12	24.33%	907
	US	CVRP1000	100	45.67	23.14%	1847
10022			200	45.04	21.43%	3816
			500	44.29	19.42%	9324
12035			0	48.24	30.05%	73
			50	45.89	23.73%	899
	CS	CVRP1000	100	45.50	22.66%	1820
			200	44.87	21.02%	3672
			500	44.07	18.82%	9180

Table 25: Comparison of uniform sampling and our coreset method with training dataset CVRP100- $\mathcal{U}(0,1)$ on test data of varying sizes. We fix the sample size as 12033.

Sample size	Method	Test distribution	RRC-budget (\downarrow)	Length (\downarrow)	$\text{Gap}\;(\downarrow)$	Time (\downarrow)
			0	856.40	9.24%	1
			50	797.45	1.72%	6
128000	Org	CVRPLib	100	797.45	1.72%	10
			200	797.45	1.72%	20
			500	797.45	1.72%	47
-			0	902.29	15.09%	1
			50	846.35	7.95%	5
	US	CVRPLib	100	838.10	6.90%	7
			200	838.10	6.90%	14
4607			500	838.10	6.90%	32
4097			0	1020.05	30.10%	1
			50	809.93	3.30%	4
	CS	CVRPLib	100	797.45	1.72%	7
			200	797.45	1.72%	14
			500	797.45	1.72%	32
			0	1096.32	39.84%	1
		CVRPLib	50	846.93	8.03%	4
	US		100	846.93	8.03%	7
			200	846.93	8.03%	13
8082			500	846.93	8.03%	32
0002			0	866.09	10.47%	1
			50	789.79	0.74%	4
	CS	CVRPLib	100	789.79	0.74%	7
			200	789.79	0.74%	13
			500	789.79	0.74%	31
			0	864.85	10.31%	1
			50	830.51	5.93%	4
	US	CVRPLib	100	797.45	1.72%	7
12033			200	797.45	1.72%	14
			500	797.45	1.72%	32
12035			0	965.89	23.20%	1
			50	816.14	4.10%	4
	CS	CVRPLib	100	809.71	3.28%	7
			200	799.16	1.93%	13
			500	787.20	0.41%	32

Table 26: Comparison of uniform sampling and our coreset method with training dataset CVRP100-U(0, 1) on test data CVRPLib.

Table 27: Comparison of uniform sampling and our coreset method with different graph embedding techniques on test data from different distributions. CS-spring is the embedding technique based on force-directed representation; CS-spectral is the spectral embedding technique; CS-MDS is the embedding technique based on multidimensional scaling.

Sample size	Method	Test distribution	Size (\uparrow)	Time (\downarrow)
		ER-[400-500]	27.40	133
4010	US	ER-[700-800]	30.36	392
		ER-[1400-1500]	34.05	1361
		ER-[400-500]	28.46	135
3973	CS-spring	ER-[700-800]	30.89	389
		ER-[1400-1500]	34.25	1361
		ER-[400-500]	27.68	132
8001	CS-spectral	ER-[700-800]	30.43	391
		ER-[1400-1500]	34.14	1362
12417		ER-[400-500]	28.43	132
	CS-MDS	ER-[700-800]	31.10	389
		ER-[1400-1500]	34.52	1361