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Trajectory Similarity Learning with Auxiliary Supervision and Optimal Matching

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Abstract

Trajectory similarity computation is a core problem in the field of trajectory data queries. However, the high time complexity of calculating the trajectory similarity has always been a bottleneck in real-world applications. Learning-based methods can map trajectories into a uniform embedding space to calculate the similarity of two trajectories with embeddings in constant time. In this paper, we propose a novel trajectory representation learning framework Traj2SimVec that performs scalable and robust trajectory similarity computation. We use a simple and fast trajectory simplification and indexing approach to obtain triplet training samples efficiently. We make the framework more robust via taking full use of the sub-trajectory similarity information as auxiliary supervision. Furthermore, the framework supports the point matching query by modeling the optimal matching relationship of trajectory points under different distance metrics. The comprehensive experiments on real-world datasets demonstrate that our model substantially outperforms all existing approaches.

1 Introduction

Evaluating the similarity between trajectories is a common task required by a wide range of applications, such as trajectory clustering [Lee *et al.*, 2007], anomaly trajectory detection [Meng *et al.*, 2019; Wu *et al.*, 2017], and road completion of maps [Chao *et al.*, 2019; Shan *et al.*, 2015]. Many trajectory distance metrics have been proposed to measure the similarity between two trajectories, including the Dynamic Time Warping (DTW) [Müller, 2007], the Fréchet distance [Eiter and Mannila, 1994], and the Hausdorff distance [Belogay *et al.*, 1997]. These metrics require quadratic computational complexity $O(l^2)$, where l is the maximum length of the trajectories. It is time-consuming when faced with long trajectories, which limits the use of these distance metrics in many real-life scenarios.

Existing trajectory similarity evaluation methods can be clustered into two categories, the classical non-learning-based methods vs. learning-based methods. Those in the former category either reduce the top- k similarity search by designing an index and pruning strategy for the trajectory database [Chen *et al.*, 2010; Shang *et al.*, 2017], or rely on an approximate solution [Driemel and Silvestri, 2017; Agarwal *et al.*, 2015]. However, they are designed by hand-crafted heuristic approaches for one or two specific distances and hence are difficult to be generalized to other distance conditions. Methods in the latter category map trajectories into a d -dimensional uniform embedding representation with learnable functions. The distance between two l -length trajectories is converted to the distance between two d -dimensional vectors. In other words, it reduces the complexity of trajectory similarity evaluation from $O(l^2)$ to $O(d)$. The improvement is considered significant as $d \ll l^2$. In addition, the representation vectors of trajectories can be used as a pre-trained embedding that could be an input of a learning-based model for downstream applications like trajectory classification. Currently, there are many trajectory representation learning methods for similarity computation [Li *et al.*, 2018; Zhang *et al.*, 2019; Chow *et al.*, 2018; Yao *et al.*, 2018; Wang *et al.*, 2019]. Among them, [Yao *et al.*, 2019] is the state-of-the-art method, which uses seed-guided neural metric learning method to compute trajectory similarity.

However, existing learning-based methods still suffer from some shortcomings. First, supervised models like [Yao *et al.*, 2019] need to calculate the distance for all trajectory pairs to construct training samples. The time complexity of this process is $O(n^2l^2)$, where n is the training set size. This cost limits the training set size of the model. In order to improve its scalability, we construct triplet training samples with a simple and fast trajectory simplification and indexing method, which can efficiently obtain approximate neighboring trajectories from massive trajectory dataset. Second, existing methods utilize the distance between two complete trajectories to evaluate their similarity but ignore the rich information carried by sub-trajectories and the mutual constraint relationship between them. We propose a novel sub-trajectory-based loss function to supervise distance between sub-trajectory pairs,

which makes our model more robust and helps to improve the performance of similarity computation. Third, a trajectory could be considered as a set of points, and distance metrics like Fréchet, DTW and Hausdorff essentially find the optimal matching between two trajectory point sets under different rules. In some scenarios, we care about these point matching relationships rather than distance values. Consequently, it is desirable to support point-matching query that locates, for a given point, its optimal matching point(s) from another trajectory. Existing methods ignore the matching relationship, so they can not support the point-matching query. We propose a point matching loss to learn the optimal matching relationships between points.

In brief, we make four contributions in this paper:

- We propose a simple yet efficient approach to locate approximate neighbors for any trajectory in a massive trajectory dataset, with the help of trajectory simplification and a tree structure index. This approach allows us to expand the size of the training set with a much lower cost and hence effectively addresses the scalability issue.
- We design a sub-trajectory distance loss function that fully leverages the distances between sub-trajectories obtained from dynamic programming for auxiliary supervision. It improves both the robustness of the model and its performance in similarity computation.
- We propose a point matching loss function to learn the optimal matching relationship between points. It allows our model to support the optimal point matching query under different distance metrics.
- We conduct comprehensive experiments on two real-world datasets to evaluate our model. The results demonstrate the advantage of our model over the state-of-the-art baselines.

2 Related Work

Existing accelerated trajectory similarity computation methods can be broadly divided into two categories, *non-learning-based* methods and *learning-based* methods.

Non-learning-based methods can be further clustered into two categories, according to the type of queries supported. The first category supports top- k nearest neighbor search. They use tree-based index structures like k-d tree, ball-tree or R-tree to maintain trajectory data hierarchically and speed up the search processing [Chen *et al.*, 2010; Shang *et al.*, 2017]. For example, [Chen *et al.*, 2010; Gowanlock and Casanova, 2015] performed pruning optimization techniques based on whether the sub-trajectories and point segments are within a bounding box, respectively. However, these methods only support top- k similarity search and can not compute the similarity between any two trajectories. The second category speeds up the computation by designing heuristic algorithms for spatial and temporal characteristics. [Driemel and Silvestri, 2017] designed a set of hash functions to speed up the computation of Fréchet and Hausdorff distance. [Chen *et al.*, 2005] proposed a method of pruning by grid histograms to approximate the similarity. [Agarwal *et al.*, 2015] proposed a fast approximate algorithm for DTW distance. These methods mainly rely on hand-crafted heuristic rules for specific

distances and hence there are room for further improvement.

Learning-based methods mainly learn trajectory embedding vectors to represent the similarity relationship through learnable functions. [Li *et al.*, 2018; Zhang *et al.*, 2019; Yao *et al.*, 2018; Wang *et al.*, 2019] all used similar encoder-decoder models to obtain trajectory vector representations and use different spatio-temporal characteristics as features. [Chow *et al.*, 2018] used the actor-critic reinforcement learning method to model trajectory distance. [Yao *et al.*, 2019] proposed a neural metric learning approach to compute the similarity with pair-wise distance as guidance. These methods neither fully mine the distance information between sub-trajectories nor explicitly restrict the matching relationship between trajectory points under different distance metrics.

3 Preliminary

Definition. A trajectory is a sequence of locations with timestamps. For any two trajectories X and Y , $S(X, Y)$ measures the similarity between them. Here, S can be the distance metrics like Fréchet distance F , DTW D , and Hausdorff distance H , etc. Our research problem is to learn the approximate similarity function $G(X, Y)$ through the neural network learning functions, such that G is close to S and can be calculated in constant time $O(d)$, where d is the dimension of the trajectory representation vector. Besides, we will also use neural network G to learn the optimal matching points in Y for points in X under different distance metrics.

Distance metrics. In this paper, we use Fréchet distance F , Hausdorff distance H , and DTW D as representative distance metrics to measure the similarity of a given trajectory pair $\langle X, Y \rangle$. They are computed differently, representing the similarity measurement of trajectories from different perspectives.

In discrete form, they can be calculated by dynamic programming [Bellman and others, 1954] or enumeration. $M_{i,j}$ ($M \in \{F, D, H\}$) refers to the corresponding distance between two sub-trajectories $X_{(1:i)}$ and $Y_{(1:j)}$. Notation $X_{(1:i)}$ refers to a sub-trajectory formed by the first i points of X . The state transition equations are as follows,

$$F_{i,j} = \max(\min(F_{i-1,j}, F_{i-1,j-1}, F_{i,j-1}), d_{p_i, q_j}) \quad (1)$$

$$D_{i,j} = \min(D_{i-1,j}, D_{i-1,j-1}, D_{i,j-1}) + d(p_i, q_j) \quad (2)$$

$$H_{i,j} = \max(\max_{t \in \{1, \dots, i\}} \min_{h \in \{1, \dots, j\}} d(p_t, q_h), \max_{t \in \{1, \dots, j\}} \min_{h \in \{1, \dots, i\}} d(q_t, p_h)) \quad (3)$$

4 Methodology

We propose a deep representation learning model Traj2SimVec as a solution. Our model consists of three parts, *training data construction*, *trajectory encoder*, and two types of *supervised loss functions*. First, *training data construction* generates triplet training samples. Second, *trajectory encoder* takes samples generated in previous step as inputs and map them to *similarity space* and *matching space* respectively to get trajectory representation vectors. Third, two types of *supervised loss functions* guide the learning in two different spaces.

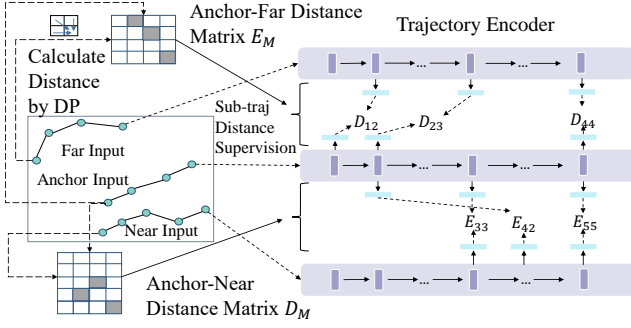


Figure 1: An example of sub-trajectory distance loss.

4.1 Training Sample Construction

Our model requires both the distance and matching relationship of trajectory pairs to supervise the learning. As reported by existing works [Faghri *et al.*, 2017], the choice of sample pairs has an essential impact on the final performance of learning models. [Yao *et al.*, 2019] enumerates the distances between any pair of trajectories to find the top- k most similar results as samples. However, it suffers from high computation cost as there are in total $O(n^2)$ trajectory pairs, making it impractical for massive datasets. For example, given a middle-sized trajectory set \mathcal{T} with $n = 100,000$ and $l = 100$, a total of $O(10^{14})$ trajectory point distance calculations are required.

We adopt a different approach by taking in similar trajectory pairs (but not necessarily the *most* similar pairs) as input to reduce the cost of forming training samples. We use a simple yet effective trajectory simplification and an index structure to obtain similar trajectory pairs as model samples quickly. Trajectory simplification [Chen *et al.*, 2009] is a compression method that reduces the size of the trajectory but retains the subject structure of the trajectory. Here, we divide the trajectory evenly into k segments and average the trajectory points of each segment as the representative point, to obtain a new trajectory of uniform length k . Given a trajectory $T = \{p_1, p_2, \dots, p_l\}$ of l points, its simplified trajectory T' is

$$T' = \left\{ \frac{1}{c_1} \sum_{i=1}^{c_1} p_i, \frac{1}{c_2} \sum_{i=c_1+1}^{c_1+c_2} p_i, \dots, \frac{1}{c_k} \sum_{i=1+l-c_k}^l p_i \right\}$$

Here, $p_i = (lat_i, lon_i)$ represents a point with latitude and longitude and $\sum_{i=1}^k c_i = l$ with $c_i \in [\lfloor \frac{l}{k} \rfloor, \lfloor \frac{l}{k} \rfloor + 1]$. A trajectory is then represented as a $2k$ -dimensional vector, in the form of $(T'.lat_1, T'.lon_1, \dots, T'.lat_k, T'.lon_k)$.

Given a trajectory set \mathcal{T} , we simplify all the trajectories in \mathcal{T} and index them using a $2k$ -dimensional k-d tree [Bentley, 1975]. Note, each point in the k-d tree corresponds to the simplified trajectory of a trajectory in \mathcal{T} . When we construct training samples, we randomly sample a point from the k-d tree as an *anchor input* I_a , and locate its k nearest neighbors with the help of k-d tree, which incurs a time complexity of $O(\log n)$. Among those k returned neighbors, we randomly select one as *near input* I_{ne} . We then randomly sample another point in the k-d tree as *far input* I_f to complete the construction of one triplet training sample, which is in the form of $\langle I_a, I_{ne}, I_f \rangle$.

4.2 Sub-trajectory Distance Loss

In order to learn the trajectory representation, we need the trajectory encoder function $f : traj \rightarrow vector$ to encode a trajectory to a vector. As shown in Figure 1, the trajectory encoder is a RNN-type model (e.g., GRU [Chung *et al.*, 2014], LSTM [Gers *et al.*, 1999] or other network structures). As introduced in Section 3, we use dynamic programming or enumeration to compute the distance between trajectories X and Y and get a distance matrix D_M . Each element D_{ij} in D_M represents the distance between sub-trajectories $X_{(1:i)}$ and $Y_{(1:j)}$. Existing methods [Yao *et al.*, 2019] only use the last element $D_{|X||Y|}$ in D_M as the supervision signal, but ignore the other crucial sub-trajectory distance constraints carried by other elements in D_M . To leverage such rich and additional supervision information, we design a sub-trajectory distance loss mechanism for auxiliary supervision.

First, we feed the original trajectories T_a , T_{ne} , and T_f in \mathcal{T} corresponding to I_a , I_{ne} and I_f in a training sample into the RNN-based trajectory encoder to obtain the corresponding hidden state vectors $H^a = \{h_1^a, \dots, h_{|T_a|}^a\}$, H^{ne} , and H^f . Then, we map the trajectories into a specific space dedicated to computing similarity through a similarity space mapping unit $f_{hid \rightarrow sim}$ that is in residual network structure. For an intermediate hidden state vector h_i , $f_{hid \rightarrow sim}$ calculates the vector v_i in the similarity space as follows.

$$C = \sigma(W_i \cdot h_i + b_i) * \tanh(W_c \cdot h_i + b_c) \quad (4)$$

$$\hat{h}_i = \sigma(W_o \cdot h_i + b_o) * \tanh(C) \quad (5)$$

$$v_i = h_i + \hat{h}_i \quad (6)$$

We can get the vector representations $V^a = \{v_1^a, \dots, v_{|T_a|}^a\}$ of trajectories T_a in the similarity space. V^{ne} and V^f can be obtained in the same way.

We can obtain the distance matrix D_M for the trajectory pair $\langle T_a, T_{ne} \rangle$ and the distance matrix E_M for the trajectory pair $\langle T_a, T_f \rangle$, based on the distance metrics calculation introduced in Section 3. Each element $D_{ij} \in D_M$ represents the real distance between the sub-trajectories $T_{a(1:i)}$ and $T_{ne(1:j)}$. The similarity between them can be modeled by the L2 distance $\|v_i^a - v_j^{ne}\|_2$. We randomly sample r elements $\{D_{i_1, j_1}, D_{i_2, j_2}, \dots, D_{i_r, j_r}\}$ from D_M as supervised signals to define the sub-trajectory loss function L_{near}^{sim} between T_a and T_{ne} . Here $D'_{i_t, j_t} = \exp(-\alpha \cdot D_{i_t, j_t}) \in [0, 1]$ is the normalized similarity of D_{i_t, j_t} , and α is a tunable parameter controlling the similarity value. We can get the sub-trajectory loss L_{far}^{sim} between T_a and T_f in a similar way.

$$L_{near}^{sim} = \frac{1}{r} \sum_{t=1}^r D'_{i_t, j_t} (D'_{i_t, j_t} - \exp(-\|v_{i_t}^a - v_{j_t}^{ne}\|_2))^2$$

In addition to the above loss function, we also add distance supervision to the entire trajectory pairs. In summary, Eq. (7) defines the sub-trajectory distance loss function, where $n_1 = |T_a|$, $n_2 = |T_{ne}|$, and $n_3 = |T_f|$.

$$L^{sim} = D'_{n_1, n_2} (D'_{n_1, n_2} - \exp(-\|v_{n_1}^a - v_{n_2}^{ne}\|_2))^2 + E'_{n_1, n_3} (E'_{n_1, n_3} - \exp(-\|v_{n_1}^a - v_{n_3}^f\|_2))^2 + L_{near}^{sim} + L_{far}^{sim} \quad (7)$$

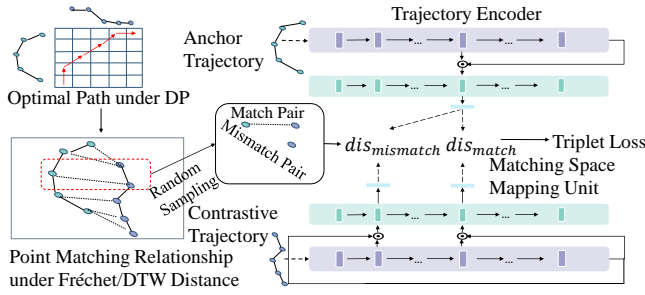


Figure 2: The example of trajectory point matching loss.

Loss function defined in Eq. (7) allows the model to learn not only the similarity between two complete trajectories, but also the similarity between different sub-trajectories. Consequently, it is equipped with the intermediate distance information of trajectory pairs. As an auxiliary supervisory loss function, it has the following two advantages. First, the intermediate distance information, to some extent, provides more trajectory pair samples and makes the model more robust. Second, the intermediate supervisory information can effectively reduce the risk of vanishing gradient of long sequences.

4.3 Trajectory Point Matching Loss

As mentioned before, the distance between two trajectories is essentially determined by the matching criteria defined by different metrics. Take Fréchet as an example. Assume an optimal path between T_a and T_{ne} is inferred by Eq. (1), denoted as $\langle (1, 1), (2, 2), (2, 3), \dots, (|T_a|, |T_{ne}|) \rangle$. This is to say, based on the optimal point matching defined by Fréchet distance, the 1st point in T_a matches the 1st point in T_{ne} , the 2nd point in T_a matches the 2nd and the 3rd points in T_{ne} , and so on. In some applications, we care about this optimal point matching relationship but not the real distance. For example, an interesting application of Fréchet distance is to indicate the shortest length of dog leash required for movement between a person and a dog. The optimal matching can tell us the corresponding positions where the dog owner took the leash when the dog was at a certain position like a garden under the Fréchet distance. The optimal matching relationship under some distance metrics (e.g., Fréchet distance and DTW) is very complicated, which requires an enumeration (e.g., dynamic programming) to get the optimal path. In order to support a faster retrieval of the optimal matching points, we propose the trajectory point matching loss, which uses the matching space mapping unit to enable the model to learn the matching relationship.

First, we construct the matching space mapping unit $f_{hid \rightarrow mat}$. Since the optimal matching is a global solution that only can be obtained after computing the entire trajectory similarity, the hidden state vector h_l needs to represent the information of the entire trajectory $T = \{p_1, p_2, \dots, p_l\}$ to help the intermediate hidden state h_t to be effectively mapped into a matching space. For an intermediate hidden state vector h_t , $f_{hid \rightarrow mat}$ uses an LSTM cell C to combine h_t and h_l in the matching space. We can get the vector representations

Dataset	Porto	Shanghai
# Trajectories (n)	999,082	1,278,550
# Trajectory points ($n \times l$)	45.6×10^6	121.7×10^6
Trajectory sampling interval	15.01 sec	9.39 sec
Trajectory minimum length	10	10
Trajectory maximum length	3836	5549
Trajectory length mean	45.67	95.19

Table 1: The description and statistics of the datasets.

M of a trajectory T in the matching space by $f_{hid \rightarrow mat}$.

$$M = \{m_1, m_2, \dots, m_l\} = f_{hid \rightarrow mat}(H) \\ = \{C_1([h_1, h_l]), C_2([h_2, h_l]), \dots, C_l([h_l, h_l])\}$$

As shown in Figure 2, for a point $p_i \in T_a$ and a point $q_j \in T_{ne}$, we learn whether they match each other based on their distance in the matching space by triplet loss [Schroff *et al.*, 2015]. We use random sampling to sample r' triples $\{\langle p_{i_1}, q_{j_1}, q_{l_1} \rangle, \dots, \langle p_{i_{r'}}, q_{j_{r'}}, q_{l_{r'}} \rangle\}$, where $\langle p_{i_t}, q_{j_t} \rangle$ s refer to the optimal matching pairs in T_a and T_{ne} and $\langle p_{i_t}, q_{l_t} \rangle$ s are the mismatching pairs. Eq. (8) defines the trajectory point matching loss between T_a and T_{ne} , where ξ is the margin value and $m_{i_t}^a$ is the matching space vector of i_t -th point p_{i_t} in T_a . We can get the trajectory point matching loss L_{far}^{match} between T_a and T_f in the same manner. Thereafter, we define the trajectory point matching loss L^{match} as $L_{far}^{match} + L_{near}^{match}$.

$$L_{near}^{match} = \frac{1}{r'} \sum_{t=1}^{r'} \min(0, \xi - \exp(-\|m_{i_t}^a - m_{j_t}^{ne}\|_2) + \exp(-\|m_{i_t}^a - m_{l_t}^{ne}\|_2)) \quad (8)$$

5 Experiments

We conduct comprehensive experiments to compare the performance of our model with existing competitors.

Datasets. We use two real trajectory datasets in our experimental study, namely *Porto* that was extracted from open source dataset available at <https://kaggle.com/c/pkdd-15-predict-taxi-service-trajectory-i> and *Shanghai*. Table 1 reports the description and statistics of the two datasets.

Hyperparameters. We split each dataset into training set, validation set and test set in the ratio 3:1:6. We set the length of simplified trajectory T' to 5, and the number of elements sampled from distance matrix D and E to 10 (i.e., $k = 5$, $r = 10$). The tunable parameter α is the reciprocal of the maximum distance of training sample distances for Fréchet and Hausdorff distance, and the mean plus three times the variance of training sample distances for DTW. The margin value ξ of point matching loss is 0.01. The sampled triple number of point matching loss is 10 (i.e., $r' = 10$). The trajectory representation vector has a dimensionality of 128 (i.e., $d = 128$). We use the LSTM model as the RNN encoder and matching space mapping unit. The hidden unit is 128. We train the model using Adam algorithm [Kingma and Ba, 2014] with an initial learning rate at 0.001. All the weights are uniformly initialized to $(-\frac{1}{\sqrt{128}}, \frac{1}{\sqrt{128}})$.

Metrics. Following the state-of-the-art approach [Yao *et al.*, 2019], we adopt HR@10, HR@50 and R10@50 as major

performance metrics. The top- k hitting ratio ($HR@k$) examines the overlap (in terms of percentage) between the returned top- k results and the ground truth; top- t recall for the top- k ground truth ($Rk@t$) evaluates the top- k ground truth recovered by the top- t produced by different methods.

Competitors. We implement *histogram* [Chen *et al.*, 2005], *traj2vec* [Yao *et al.*, 2018], *t2vec* [Li *et al.*, 2018], *NT-No-SAM* [Yao *et al.*, 2019], and *NEUTRAJ* [Yao *et al.*, 2019] as representatives. *histogram* is a non-learning based embedding method, which splits a region into grids and approximates the similarity by counting the number of trajectory points in different grids. *traj2vec* and *t2vec* are both auto-encoder models, but *t2vec* uses additional embedding and attention techniques and processes the data noise. *NEUTRAJ* is the state-of-the-art trajectory similarity learning method with its source code openly available at <https://github.com/yaodi833/NeuTraj>. It is a metric learning approach using seed-guided neural network to compute similarity. *NT-No-SAM* is a weaker version of *NEUTRAJ*, which replaces the spatial attention module with standard LSTM.

5.1 Overall Evaluation

The performance of similarity search. We evaluate the performance of similarity computation approaches for the top- k similarity search task. We randomly selected 10,000 trajectories from the test set, and try to find top- k similar trajectories from the entire test set (600k+ and 700k+ trajectories in Porto and Shanghai) for each of these trajectories. Since *NEUTRAJ* and *NT-No-SAM* need to calculate the distance matrix for all trajectory pairs in the training set to construct training samples, they cannot handle large datasets. Instead, we randomly select 10,000 trajectories from the training set to form their training samples. As reported in Table 2, our model *Traj2SimVec* outperforms all the competitors substantially. Compared with state-of-the-art algorithm (*NEUTRAJ*), *Traj2SimVec* performs much better. For example, it achieves at least 25%+ gain in terms of $HR@10$ on both datasets. Although *Traj2SimVec* and *NEUTRAJ* both use metric learning methods to learn the similarity of trajectories, *Traj2SimVec* performs better since it is scalable to large-scale datasets and it uses a sub-trajectory distance loss for extra auxiliary supervision. In addition, we report the testing time, preprocessing and training time required by different methods in Figure 3. We compare the non-vectorized method to directly query the similarity of original trajectories with learning models. The results show that the testing time for learning models is significantly shorter than non-vectorized method. We could also observe that *NEUTRAJ* and *Nt-No-SAM* require significantly longer preprocessing time than other methods, as they have to calculate the trajectory distance matrix for training.

The performance of point matching. We evaluate the performance of point matching approaches for the top- k matching task. We implement two baselines, namely *Nearest* and *Order*. *Nearest* finds the nearest neighbor in the corresponding trajectory as a matching point for every query point. *Order* uses the relative position of each point in the entire trajectory as a distance metrics. Given a trajectory $T_1 = \{p_1, \dots, p_5\}$, the relative position of point $p_3 \in T_1$ is $3/5$.

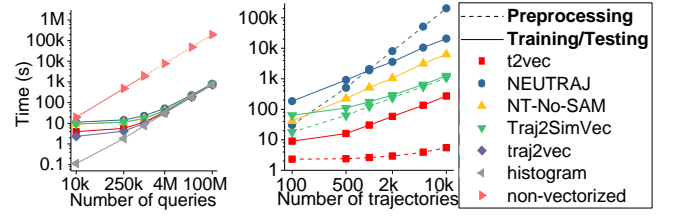


Figure 3: The testing time (left), preprocessing and training time (right) of different methods on Fréchet distance under Porto dataset.

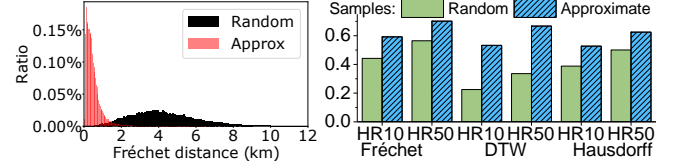


Figure 4: Distance distribution of training samples and performance of *Traj2SimVec* under different sampling approaches.

Consequently, its nearest point in another trajectory T_2 is the one located at the same relative position. The point optimal matching relationship in Hausdorff distance finds the point in another trajectory that is closest to it, which is equivalent to the baseline *Nearest*. However, the point optimal matching relationship under the distance metrics of DTW and Fréchet is much more complicated. We use $M@1$, $M@5$ and $M@10$ as performance metrics. $M@k$ examines the overlap percentage of the returned top- k nearest points and the ground truth optimal matching points. As shown in Table 3, *Traj2SimVec* consistently outperforms both baseline methods with noticeable advantage. It shows that the distance between points and the order of points in the trajectory cannot characterize the relationship of point optimal matching well. Our model learns more complex matching rules than baselines.

5.2 Performance of *Traj2SimVec*

The effectiveness of approximate sampling. Figure 4 reports the distribution of distances of trajectory pairs in training set under two different sampling approaches. Random sampling (RS) randomly selects a trajectory from the training set to form a sample pair with anchor trajectory T_a . Approximate nearest sampling (AS) uses trajectory simplification and indexing to find a trajectory close to T_a to form a trajectory pair with T_a . As reported, samples formed by AS have much shorter distances, as compared with those generated by RS. This demonstrates the superior performance of the training sample construction component of *Traj2SimVec*. We also report the performance of our model trained by different training samples under Porto dataset in Figure 4. As observed, samples generated by AS approach are more powerful as they help the learning model to achieve a much better performance. That is to say, our model can construct approximate near and far samples to effectively train on a large-scale dataset, which enhances the scalability of our model.

The effectiveness of sub-trajectory distance loss. In order to demonstrate the effectiveness of the proposed sub-trajectory distance loss with auxiliary supervision, we com-

Dataset	Method	Fréchet			DTW			Hausdorff		
		HR@10	HR@50	R10@50	HR@10	HR@50	R10@50	HR@10	HR@50	R10@50
Porto	histogram	0.0111	0.0255	0.0336	0.0149	0.0331	0.0420	0.0115	0.0269	0.0359
	traj2vec	0.1039	0.1457	0.3003	0.1336	0.1789	0.3729	0.0931	0.1236	0.2598
	t2vec	0.2401	0.2832	0.4655	0.2476	0.2922	0.4819	0.2219	0.2543	0.4313
	NT-No-SAM	0.4727	0.5909	0.8021	0.3931	0.4979	0.7329	0.4024	0.5004	0.7512
	NEUTRAJ	0.4751	0.5928	0.8071	0.4126	0.5223	0.7525	0.3992	0.5029	0.7484
	Traj2SimVec	0.5919	0.7017	0.9011	0.5332	0.6669	0.8988	0.5270	0.6251	0.8694
Shanghai	histogram	0.0212	0.0393	0.0547	0.0270	0.0488	0.0666	0.0238	0.0463	0.0657
	traj2vec	0.0936	0.1317	0.2888	0.1047	0.1413	0.3199	0.0667	0.0966	0.2229
	t2vec	0.2735	0.2921	0.5021	0.2690	0.2856	0.4849	0.2064	0.2202	0.4234
	NT-No-SAM	0.4663	0.5894	0.7628	0.2484	0.3393	0.5674	0.3260	0.3999	0.7077
	NEUTRAJ	0.4774	0.5981	0.7944	0.2597	0.3497	0.5857	0.3272	0.4032	0.7092
	Traj2SimVec	0.6136	0.6910	0.8902	0.3626	0.4829	0.7376	0.4805	0.5625	0.8867

Table 2: The performance of similarity computation approaches on Fréchet, Hausdorff and DTW distances under Porto and Shanghai dataset.

Dataset	Porto (Fréchet)			Porto (DTW)			Shanghai (Fréchet)			Shanghai (DTW)		
Method	M@1	M@5	M@10	M@1	M@5	M@10	M@1	M@5	M@10	M@1	M@5	M@10
Nearest	0.1484	0.2995	0.4361	0.1931	0.3318	0.4482	0.2125	0.2991	0.3794	0.2265	0.3047	0.3729
Order	0.1087	0.2979	0.4688	0.1470	0.4147	0.6127	0.0563	0.1548	0.2585	0.0723	0.2066	0.3368
Traj2SimVec	0.3353	0.6649	0.8010	0.3338	0.7373	0.8780	0.3153	0.5602	0.6872	0.3049	0.5442	0.6799

Table 3: The performance of point optimal matching approaches on Fréchet and DTW distances under Porto and Shanghai dataset.

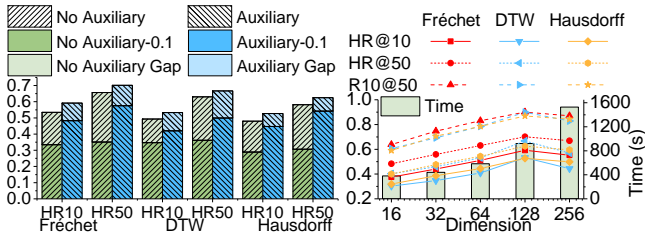


Figure 5: Left: The performance of Traj2SimVec with and without sub-trajectory distance loss as auxiliary supervision. Right: The performance under different vector dimensions.

pare it with the model without the auxiliary supervision. As shown in Figure 5, the model using sub-trajectory distance loss as auxiliary supervision performs better than the model without sub-trajectory distance loss. We also keep only the first 10% sub-trajectory for each trajectory in the test set, and conduct experiments on the sub-trajectories to test the robustness of the model, denoted as *Auxiliary-0.1* and *No Auxiliary-0.1* in the left subfigure of Figure 5. We could observe from the result that the gap of decline in the model with auxiliary supervision is significantly smaller than the model without auxiliary supervision. This proves that our sub-trajectory distance loss not only improves the overall performance, but also effectively supervises the similarity relationship of sub-trajectories, making the model more robust.

The impact of the dimensions of representation vector. In all the above experiments, a trajectory is represented by a 128-dimensional vector. In order to evaluate the impact of the dimensions d of the vector, we evaluate the performance Traj2SimVec under different settings (i.e., 16, 32, 64, 128, and 256), with result reported in the right subfigure of Figure 5. It is observed that Traj2SimVec gradually improves its performance as the vector dimension increases, and it reaches

its best performance when $d = 128$. This shows that the representation vector of a trajectory needs enough large dimension to represent the spatio-temporal information of the trajectory, but a space of too many dimensions actually increases the difficulty of learning. In addition, we report the time cost of computing vector distance and performing top- k similarity search on the entire test set under different d settings. We test the time cost on an AMD Ryzen Threadripper 2950X processor with 32 threads. As d increases, it only requires more time for computing the trajectory similarity. However, as compared with computing similarity between original trajectories that requires more than 20 hours, the evaluation of trajectory similarity in this vector space is far more efficient.

6 Conclusion

In this paper, we propose a novel trajectory similarity learning model Traj2SimVec. It adopts trajectory simplification and indexing technique to reduce the cost of generating samples, hence making it applicable to large datasets; it proposes a sub-trajectory distance loss to further improve the robustness and performance of the model; and it utilizes a point matching loss to learn the optimal matching between points.

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