Computing Trajectory Similarity in Linear Time: A Generic Seed-Guided Neural Metric Learning Approach

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Abstract—Trajectory similarity computation is a fundamental problem for various applications in trajectory data analysis. However, the high computation cost of existing trajectory similarity measures has become the key bottleneck for trajectory analysis at scale. While there have been many research efforts for reducing the complexity, they are specific to one similarity measure and often yield limited speedups. We propose NEUTRAJ to accelerate trajectory similarity computation. NEUTRAJ is generic to accommodate any existing trajectory measure and fast to compute the similarity of a given trajectory pair in linear time. Furthermore, NEUTRAJ is elastic to collaborate with all spatialbased trajectory indexing methods to reduce the search space. NEUTRAJ samples a number of seed trajectories from the given database, and then uses their pair-wise similarities as guidance to approximate the similarity function with a neural metric learning framework. NEUTRAJ features two novel modules to achieve accurate approximation of the similarity function: (1) a spatial attention memory module that augments existing recurrent neural networks for trajectory encoding; and (2) a distance-weighted ranking loss that effectively transcribes information from the seed-based guidance. With these two modules, NEUTRAJ can yield high accuracies and fast convergence rates even if the training data is small. Our experiments on two real-life datasets show that NEUTRAJ achieves over 80% accuracy on Fréchet, Hausdorff, ERP and DTW measures, which outperforms stateof-the-art baselines consistently and significantly. It obtains 50x-1000x speedup over bruteforce methods and 3x-500x speedup over existing approximate algorithms, while yielding more accurate approximations of the similarity functions.

Index Terms—deep metric learning, trajectory similarity, linear time

I. INTRODUCTION

Computing the similarity between two trajectories is a primitive that is fundamental to many searching and mining tasks for trajectory analysis. Various measures have been proposed to capture the intrinsic structural similarities between trajectories, including Dynamic Time Warping(DTW) [31], the Hausdorff distance [3], the Fréchet distance [2], Edit distance with Real Penalty(ERP) [9], etc. These similarity measures have played a key role in tasks like anomaly detection [18], duplicate detection [27], trajectory clustering [6].

Unfortunately, the high computation cost of existing trajectory similarity measures has become the *de facto* bottleneck

for trajectory analysis at scale. To compute the similarity between two trajectories, existing techniques often require to align the points in the two trajectories, accumulate the information among all aligned pairs, and finally produce the distance. Such a process incurs quadratic and even superquadratic time complexity and limits many trajectory mining algorithms to scale to large datasets. For instance, it takes us more than 6.5 hours to compute the pair-wise Hausdorff distances for merely $\sim\!8000$ human GPS trajectories on a highend server. As massive trajectory data are being collected at an unprecedentedly massive scale in many kinds of scenarios, fast trajectory similarity computation under different measures has become a pressing need.

The difficulties in fast trajectory similarity computation are two-fold. The first is the complicated nature of trajectory similarity computation. A pair of input trajectories may have completely different lengths, and the best alignment of the two trajectories is subject to flexible shifting and scaling instead of exact head-to-tail matching. As such, most existing techniques have to employ a scan-and-align mechanism for determining the best matching, and it is hard to decouple the matching process and reduce the time cost. The second is the variations across different similarity measures. Prevailing trajectory measures (e.g., DTW, Hausdorff, Fréchet) differ a lot in their definitions and computation mechanisms. It is challenging to design a generic accelerating strategy that accommodates all the existing measures.

There have been considerable research efforts attempting to accelerate trajectory similarity computation for various tasks. Such efforts can be generally categorized into two lines. The first [10], [14], [15], [30] is to reduce the involved number of computations at a global level. However, the techniques along this line are exclusively designed for the top-k similarity search task. Instead of reducing the computation complexity for an ad-hoc pair of trajectories, they focus on designing indexing and pruning strategies for a given trajectory database and reducing the number of computations for top-k similarity search. As such, they cannot be applied for tasks that require the distances between all trajectory pairs such as trajectory



clustering and anomaly detection. The second line aims at directly reducing the time complexity for trajectory similarity computation with approximate algorithms. Different strategies have been proposed for different measures, such as locality sensitive hashing (LSH) for the Fréchet and DTW distances [12]. Unfortunately, the techniques are designed for one specific distance measure and not applicable to any other measure.

We propose a model that drastically accelerates trajectory similarity computation for any measures. Our proposed model, named NEUTRAJ, is an approximate approach based on neural metric learning. NEUTRAJ samples a pool of seed trajectories from the database and use their pair-wise similarities as guidance. Specifically, NEUTRAJ learns a neural network that jointly embeds input trajectories and approximates the distance function. It has the following attractive characteristics:

- Generic: Unlike previous methods that are specific to one trajectory similarity measure, NEUTRAJ is generic enough to support any existing measure. It can thus be used for accelerating most trajectory mining tasks.
- Fast: Given an ad-hoc pair of trajectories, NEUTRAJ is able to compute their similarity in O(L) time complexity, where L is the length sum of the pair. In practice, we observed it at least 50x faster than accurate bruteforce computation.
- Accurate: NEUTRAJ achieves superb approximation performance in practice. On two real trajectory datasets, NEUTRAJ achieves over 80% hitting ratio and less than 50m average error distance on top-10 similar trajectory search task for Fréchet, Hausdorff, DTW and ERP. Meanwhile, it obtains more than two times higher hitting ratios compared with state-of-the-art trajectory similarity approximation methods [12].
- Elastic: NEUTRAJ embeds the trajectory without losing spatial information which makes it elastic to extend by other indexing and pruning strategies. In tasks that similarities of all pairs are non-essential, NEUTRAJ is able to cooperate with existing indexing methods [10], [15], [30] for reducing the computing space.

The core of NEUTRAJ is a deep metric learning framework that uses recurrent neural networks (RNNs) to generate trajectory embeddings. We sample a pool of seed trajectories from the database and compute their pair-wise similarities. With the computed seed similarities as guidance, we design a pairwise loss to optimize the network for fitting seed similarities. NEUTRAJ features two novel modules that encourage the network to approximate the similarity function accurately: (1) Spatial attention memory (SAM). Vanilla RNNs along with its existing variants (GRU, LSTM) can only model one sequence without considering between-sequence correlations. Our designed SAM unit memorizes the information from previously processed trajectories with the attention mechanism, and capture the correlations between training trajectories to produce better trajectory embeddings. (2) Distance-weighted ranking loss. One difficulty of making use of the seed trajectories is the dilemma between efficiency and effectiveness.

On the one hand, training the network sufficiently would preferably iterate over all pairs of trajectories. On the other hand, a full enumeration of all pairs of trajectories incurs expensive computation time. To address this dilemma, we propose a distance-weighted sampling strategy to focus on the more discriminative training pairs. Along with the weighted sampling strategy, distance-weighted ranking loss is a ranking loss that learns the parameters of the network to conform to the guidance from the seeds. With these two novel modules, NEUTRAJ can yield high accuracies and fast convergence rates even if the training data is small.

Our contributions can be summarized as follows:

- We propose a neural metric learning method for accelerating trajectory similarity computation under different measures. To the best of our knowledge, NEUTRAJ is the first method that supports accelerating generic trajectory similarity measures, making it widely applicable to many applications.
- We propose the spatial attention memory unit to model the correlation between spatially close trajectories based on an attention network and external memory tensor.
- 3) We design a weighted sampling and learning module that fully unleashes the power of seed trajectories. Compared with existing architecture, our learning module yields faster convergence rates and higher accuracies.
- 4) We conduct extensive experiments on two real trajectory datasets and four popular trajectory similarity measures. The results demonstrate that the proposed model consistently outperforms state-of-the-art baselines in both accuracy and efficiency.

II. RELATED WORK

In this section, we provide an overview of existing studies related to NEUTRAJ from three perspectives: (1) trajectory similarity computation; (2) deep metric learning; and (3) memory networks.

Trajectory Similarity Computation. Various techniques have been proposed to accelerate trajectory similarity computation, which can be broadly categorized into two categories. The first category uses indexing and pruning techniques to reduce the involved number of computations at a global level. Most techniques in this category employ tree-based index structures [10], [11], [14], [19], [27], such as K-D tree or R-tree to organize the trajectory data in a hierarchy. Based on the index, bounding-box-based pruning techniques are employed to eliminate unnecessary computations. Thus sub-trajectories [10], [11] or point segments [15], [30] in a bounding box which are too faraway to belong to the top-k results are pruned. However, the techniques in this category are specifically designed for the top-k similarity search problem. They do not reduce the time complexity of computing the similarity between a pair of trajectories. Hence, they cannot be applied for tasks that require the distances of all pairs such as trajectory clustering and anomaly detection.

The second category aims at designing approximate algorithms to speed up similarity computation for a pair of trajec-

tories. Most techniques in this category treat each trajectory as a spatial curve and address the problem from the angel of computational geometry. Focusing on the Hausdorff distance, Farach-Colton et al. and Backurs et al. [4], [13] proposed embedding-based methods for approximating nearest neighbor search. Salvador et al. [1] proposed an approximate algorithm which can fast compute the DTW distance. Thanawin et al. [26] proposed a method which omits the square computation step to speed up DTW computation. Li et al. [20] proposed a new trajectory similarity measure based on road network and employ deep representation learning to approximate it. Very recently, Driemel et al. [12] proposed a locality sensitive hashing (LSH) based algorithm for fast computing the Fréchet and Hausdorff distances. Although these algorithms can achieve high computation efficiency, they suffer from two shortcomings. First, they rely on hand-crafted heuristics and could lead to unsatisfactory accuracies. In our experiments, we observed that these algorithms generate poor approximations in many cases. Second, they are all designed for one or two specific measures. It is hard to adapt these techniques for other similarity measures.

Deep Metric Learning. NEUTRAJ is related to the recent development of deep metric learning, which aims at learning a distance function that measures how similar two objects are based on neural networks. Bromley *et al.* [5] pioneered deep metric learning and proposed the classic Siamese network for signature verification. Qian *et al.* [25] used precomputed activation features to learn a feature embedding for classification. Pei *et al.* [24] extended the Siamese network to learn a similarity metric for sentences. Our method differs from the above models in two aspects. First, they are all designed for modeling one sequence independently, while ours employs the spatial attention mechanism to capture the correlations among all the trajectories. Second, they all use random sampling to generate training samples and could suffer from low convergence for trajectory data.

Memory Network. NEUTRAJ employs memory network to capture the relations between trajectories. The embryonic form of memory network is proposed to solve the tasks that need to model long term dependency [16]. Weston *et al.* [29] first employed a long-term memory component and defined the read and write operation for questing answering(QA). Then Sukhbaatar *et al.* [28] extended the architecture to recurrent neural network. In the proposals [8], [29], the memory network was extended to a hierarchical structure. But these memory structures are designed without considering the locality information and cannot be used directly for trajectory modeling.

III. PRELIMINARIES

A. Problem Definition

We consider a trajectory database $\mathscr T$ and a trajectory similarity function $f(\cdot,\cdot)$. Each trajectory $T\in\mathscr T$ is a sequence of points recording the trace of a moving object. Although each sample point in a trajectory has a sampling time, we only focus on finding trajectories with similar

TABLE I NOTATIONS USED IN THIS PAPER

Notations	Description
\mathscr{T},\mathfrak{S}	A trajectory database and a pool of N seeds trajectories
_	which are random sampled from the database.
T	A trajectory in \mathcal{T} which consist of a sequence of
\mathbf{D}, \mathbf{S}	coordinate tuples. The distance and similarity matrices of $\mathfrak S$ which have
\mathbf{D}, \mathbf{S}	the same size: $\mathbb{R}^{N \times N}$.
\mathbf{M}	The memory tensor which stores the spatial informa-
	tion of $P \times Q$ grids.
$\mathbf{E_i}, \mathbf{E_j}$	d-dimensional embedding vectors of T_i and T_j learnt
	by NeuTraj.
X_t	The input of NEUTRAJ at t-time step which contains
W, U, b	the coordinate input X_t^c and grid input X_t^g . The linear weights and bias in SAM unit.
$\mathbf{f_t}, \mathbf{i_t}, \mathbf{o_t}$	The forget, input and output gates in SAM-argumentd
it, it, Ot	LSTM unit.
$\mathbf{s_t}$	Novel spatial gate in SAM unit which controls the
	read and write operations on M.
$\mathbf{c_t}, \mathbf{\hat{c_t}}$	The cell state and intermediate cell state in SAM
	unit at t-time step which store the information of the
1. 1.	processed $t-1$ steps.
$egin{array}{c} \mathbf{h_t}, \mathbf{h_{t-1}} \ \mathbf{G_t} \end{array}$	The hidden states in SAM unit at time step t and $t-1$. The spatial information matrix of X_t with shape
G_{t}	$\mathbb{R}^{(2w+1)^2 \times d}$, where w is the scanning bandwidth.
Α	The spatial attention weight that reflects the similarity
11	weights of $\hat{\mathbf{c}}_t$ over \mathbf{G}_t .
$\mathbf{c_t^{cat}}, \mathbf{c_t^{his}}$	The intermediate concatenated state and the final his-
- , -	torical state in memory $read$ operation at t -time step.
T_a	The anchor trajectory which is sampled from the seeds
- a - d	for training NEUTRAJ.
$\mathscr{T}_a^s, \mathscr{T}_a^d$	n similar trajectories and n dissimilar trajectories of
	T_a which are sampled from the seeds for fitting the
$\mathbf{S}_a^s, \mathbf{S}_a^d$	pair-wise similarities. The ground truth similarities of both similar and dis-
$\mathfrak{S}_a,\mathfrak{S}_a$	similar pairs of T_a
$\hat{\mathbf{S}_a^s}, \hat{\mathbf{S}_a^d}$	The similarities of T_a which are calculated by NEU-
$\mathfrak{S}_a,\mathfrak{S}_a$	TRAJ.

shape, regardless of the time information. Without loss of generality, we consider two-dimensional trajectories. That is, each trajectory $T = [X_1^c, ..., X_t^c, ...]$ is a sequence of tuples where $X_t^c(x_t, y_t)$ is the t-th location of the object. For any two trajectories $T_i, T_j \in \mathcal{T}$, function $f(T_i, T_j)$ measures the similarity between T_i and T_j . Here, $f(\cdot, \cdot)$ could be the DTW similarity, the Hausdorff distance, the Fréchet distance, or any other trajectory similarity measure. We omit the detailed definitions of these measures due to the space limit.

Our problem is to compute the similarity for an ad-hoc pair of trajectories from $\mathscr T$ under the similarity function $f(\cdot,\cdot)$. However, for most prevailing similarity measures, computing the similarity between a pair of trajectories incurs quadratic time complexity. Hence, the research question is: how can we learn an approximate similarity function $g(\cdot,\cdot)$, such that computing $g(T_i,T_j)$ takes O(n) time while the difference $|f(T_i,T_j)-g(T_i,T_j)|$ is minimized.

B. Overview of NEUTRAJ

At the high level, NEUTRAJ adopts a neural metric learning framework. It randomly samples N trajectories from $\mathcal T$ as the

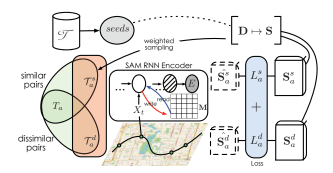


Fig. 1. Architecture of NeuTraj. Taking similar and dissimilar pairs of anchor T_{a_i} as input, NeuTraj first generates the embedding of each trajectory, and then fits the pair-wise similarity guided by the ground truth in S.

pool of seeds $\mathfrak S$ and computes a symmetric $N\times N$ distance matrix $\mathbf D$ for $\mathfrak S$. Then it transforms the original distance matrix into a normalized similarity matrix $\mathbf S$. Leveraging the matrix $\mathbf S$ as guidance, NEUTRAJ further learns a neural network, which maps arbitrary-length trajectories into low-dimensional space to capture their similarities. More formally, for any two input trajectories T_i and T_j $(i,j\in[1,...,N])$, NEUTRAJ projects them to two d-dimensional vectors $\mathbf E_i$ and $\mathbf E_j$, respectively. The learned mapping should be similarity preserving, namely $f(T_i,T_j)\approx g(T_i,T_j)$ where $g(\cdot,\cdot)$ is the similarity between $\mathbf E_i$ and $\mathbf E_j$ in the embedding space. Figure 1 illustrates the architecture of NEUTRAJ. It consists of two major parts: spatial attention memory(SAM) augmented RNN encoder and seed-guided metric learning method.

SAM Augmented RNN Encoder. NEUTRAJ relies on recurrent neural networks (RNN) to model the trajectory and takes the last hidden state of RNN as the embedding vector. However, as aforementioned, vanilla RNNs and its variants (GRU, LSTM) capture the information of each sequence independently. For trajectory similarity computing, the correlations between trajectories are critical. It is important to leverage the information of spatially close trajectories previously seen to guide the metric learning process. Thus, we design a spatial attention memory module in NEUTRAJ. It employs a spatial memory tensor to store the spatial information of previously processed trajectories. The memory tensor underpins *read* and *write* operations over the entire space based on the soft attention mechanism, such that the information of previously seen trajectories can be encoded and retrieved on demand.

Seed-Guided Neural Metric Learning. Based on SAM augmented RNN, NEUTRAJ builds a seed-guided metric learning architecture to consume a pair of trajectories, and learns the network to approximate the similarity matrix **S**. Existing metric learning methods employ random sampling to produce training pairs, which implies all trajectories are equally weighted. But this assumption dose not hold in trajectory metric learning as it ignores the spatial proximity between trajectories. Particularly, we develop a distance-weighted sampling procedure and a ranking loss objective to solve this problem. Unlike previous random sampling, the distance-

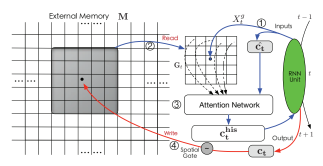


Fig. 2. Illustration of the proposed spatial attention memory(SAM) with scanning bandwidth w=2. At each time step, SAM takes two inputs, the input grid cell X_t^g and the intermediate cell state $\hat{\mathbf{c_t}}$. The reader first scans the memory \mathbf{M} to get $(2*2+1)^2=25$ grid cell embeddings. Then it calculates and outputs the attention cell state $\mathbf{c_t^{his}}$. The writer (red lines) updates \mathbf{M} with the cell state $\mathbf{c_t}$ in the recurrent unit.

weighted sampling focuses on the more discriminative training pairs from the seed trajectories. With the weighted sampling strategy, each seed trajectory T_a is associated with one similar list \mathscr{T}_a^s and one dissimilar list \mathscr{T}_a^d in the pool. The ranking loss then learns the parameters of the network for fitting the similarities to \mathbf{S} and preserving the ranking order in both similar and dissimilar pairs.

IV. SAM AUGMENTED RNN ENCODER

In this section, we introduce the Spatial Attention Memory (SAM) module that augments existing RNN architectures for trajectory encoding. Below, we first introduce the spatial attention memory structure. Then we present a fancy RNN unit, SAM-augmented LSTM, which augment existing recurrent neural networks with the SAM. Finally, we detail the *read* and *write* operations of the memory in SAM-augmented LSTM.

A. Grid-Based Memory Tensor

The SAM module is a grid-based memory network. As a prepossessing step, we partition the space into small grid cells. Then any trajectory $T = [X_1^c, ..., X_t^c, ...]$ can be mapped into a sequence $T^g = [X_1^g,...,X_t^g,...]$ where $X_t^g = (x_t^g,y_t^g)$ specifies the grid cell at t - th position. Figure 2 shows the architecture of proposed SAM module. As shown, the core part of SAM is a memory tensor M. The memory tensor M stores vector representations for all the grid cells in the space, which enable encoding and retrieving information for previously seen trajectories. Formally, assume the entire space is partitioned into $P \times Q$ grid cells, then the dimensionality of the memory tensor is $\mathbb{R}^{P \times Q \times d}$, where d is the hidden size of the recurrent unit. Each slice (p, q, :) in M stores the embedding vector of cell (p,q) and all grid cell embeddings are initialized with 0 before training. As NEUTRAJ continuously processes the trajectories in the training data, the memory tensor M will be updated accordingly by memory-augmented recurrent unit to encode the information in processed trajectories.

B. Memory-Augmented RNN Unit

The SAM module allows for memorizing and retrieving information from processed trajectories. We leverage it to

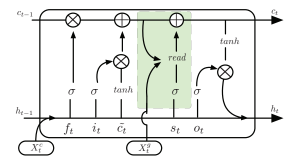


Fig. 3. SAM-augmented LSTM.

ameliorate standard RNN unit. In this way, the RNN encoder captures the information from not only the current trajectory, but also those similar ones previously processed. In what follows, we show the details of SAM-augmented LSTM.

Figure 3 shows the architecture of SAM-augmented LSTM unit and the green parts are the novel SAM module. As shown, at each time step t, the unit takes $X_t = (X_t^c, X_t^g)$ and the previous hidden state $\mathbf{h_{t-1}}$ as the input and outputs $\mathbf{h_t}$ to the next recurrent step. As in LSTM, SAM-augmented LSTM employs gating mechanism to control the operations on the cell state $\mathbf{c_t}$ that stores core information of the processed sequence. The recurrent step is performed as follows:

$$(\mathbf{f_t}, \mathbf{i_t}, \mathbf{s_t}, \mathbf{o_t})^T = \sigma(\mathbf{W_g} \cdot X_t^c + \mathbf{U_g} \cdot \mathbf{h_{t-1}} + \mathbf{b_g})$$
 (1)

$$\tilde{\mathbf{c_t}} = \tanh(\mathbf{W_c} \cdot \mathbf{X_t^c} + \mathbf{U_c} \cdot \mathbf{h_{t-1}} + \mathbf{b_c})$$
 (2)

$$\hat{\mathbf{c_t}} = \mathbf{f_t} \cdot \mathbf{c_{t-1}} + \mathbf{i_t} \cdot \tilde{\mathbf{c_t}} \tag{3}$$

$$\mathbf{c_t} = \hat{\mathbf{c_t}} + \mathbf{s_t} \cdot read(\hat{\mathbf{c_t}}, X_t^g, \mathbf{M}) \tag{4}$$

$$write(\mathbf{c_t}, \mathbf{s_t}, X_t^g, \mathbf{M})$$
 (5)

$$\mathbf{h_t} = \mathbf{o_t} \cdot \tanh(\mathbf{c_t}) \tag{6}$$

where $\mathbf{W_g} \in \mathbb{R}^{4d \times 2}$, $\mathbf{U_g} \in \mathbb{R}^{4d \times d}$, $\mathbf{W_c} \in \mathbb{R}^{d \times 2}$, $\mathbf{U_c} \in \mathbb{R}^{d \times d}$ and d is the hidden state size. All of the gates($\mathbf{f_t}, \mathbf{i_t}, \mathbf{s_t}, \mathbf{o_t}$), cell states($\tilde{\mathbf{c_t}}, \hat{\mathbf{c_t}}, \mathbf{c_t}$) and hidden states($\mathbf{h_t}, \mathbf{h_{t-1}}$) have the same shape: $\mathbb{R}^{d \times 1}$.

To obtain the hidden state $\mathbf{h_t}$, the unit performs following steps: (1) Gate operations. By Equation 1, the unit applies a sigmoid function σ on the linear transformation of the coordinate input X_t^c and the previous hidden state $\mathbf{h_{t-1}}$ to obtain the four gates: forget gate $\mathbf{f_t}$, input gate $\mathbf{i_t}$, spatial gate $\mathbf{s_t}$ and output gate $\mathbf{o_t}$. (2) Cell state operations. By Equations $2 \sim 4$, the unit produces cell state of current time-step $\mathbf{c_t}$, based on the $\mathbf{f_t}$, $\mathbf{i_t}$, $\mathbf{s_t}$ and the inputs (X_t^c, X_t^g) . (3) Hidden state operations. By Equation 6, the unit generates $\mathbf{h_t}$ and output it to the next recurrent step.

The main novelty of SAM-augmented LSTM lies in Equation 4 that augments $\hat{c_t}$ with historical information by the read operations. After that, the unit updates the memory tensor M by Equation 5. The details of read and write operations are described in the next section.

C. Attention-Based Reads and Writes

With the memory tensor M, NEUTRAJ uses the attention mechanism to capture the information in processed trajectories: (1) the *read* operation retrieves relevant grid cell

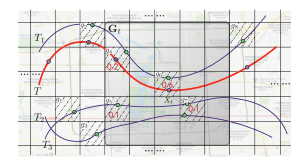


Fig. 4. Example to illustrate spatial reads and writes, $T_1 \sim T_3$ are previously processed trajectories; T is the current processing trajectory and X_t is the current processing input. After processed $T_1 \sim T_3$, grid cell embeddings of $g_1 \sim g_8$ are updated to non-zero value. By spatial readers, NEUTRAJ scans the spatial closets grid cells of X_t and computes the attention weights of grids for encoding T. The attention weights of non-zero grid cells $g_4 \sim g_7$ are the red values. After that, the cell state of X_t is used to update the grid cell embedding of g_6 .

embeddings from the M to augment encoding the current trajectory; and (2) the *write* operation attends to relevant grid cells and updates grid cell embeddings with the information of the current trajectory.

1) Spatial Memory Reader: The reader retrieves information from the memory and uses the information to augment RNN-based trajectory encoding. As shown in Figure 2-(1), at each step t, the reader takes two inputs: (1) grid cell input X_t^g ; and (2) intermediate cell state $\hat{\mathbf{c_t}}$. With these two inputs, the reader outputs a vector $\mathbf{c_t^{his}}$, which augments $\hat{\mathbf{c_t}}$ with the influence of previously processed trajectories close to X_t^g .

Illustrated by Figure 2-②, the attentional reader first looks up the grid cells that are spatially close to $X_t^g = (x_t^g, y_t^g)$. Specifically, the reader uses the bandwidth w to perform a memory scan and identifies the grid cells around (x_t^g, y_t^g) : $scan(x_t^g) = [x_t^g - w, x_t^g + w]$; $scan(y_t^g) = [y_t^g - w, y_t^g + w]$. The read grid cell embeddings are stored in a matrix \mathbf{G}_t with shape $\mathbb{R}^{(2w+1)^2 \times d}$. After memory scan, the reader employs the attention network to transform the matrix \mathbf{G}_t to an d-dimensional vector, which is shown in Figure 2-③. The attention mechanism is performed as follows:

$$\begin{split} \mathbf{A} &= \operatorname{softmax}(\mathbf{G_t} \cdot \hat{\mathbf{c_t}}); \quad mi\mathbf{x} = \mathbf{G_t}^{\mathrm{T}} \cdot \mathbf{A}; \\ \mathbf{c_t^{cat}} &= [\hat{\mathbf{c_t}}, mi\mathbf{x}]; \quad \mathbf{c_t^{his}} = \tanh(\mathbf{W_{his}} \cdot \mathbf{c_t^{cat}} + \mathbf{b_{his}}) \end{split}$$

where the matrix $\mathbf{W_{his}}$ and $\mathbf{b_{his}}$ are the parameters of the attention network. $\mathbf{A} \in \mathbb{R}^{(2w+1)^2 \times 1}$ is the attention weight that reflects the similarity of the $\hat{\mathbf{c_t}}$ over the historical grid cell embedding matrix $\mathbf{G_t}$. For example, Non-zero attention weights over grid cells $g_4 \sim g_7$ in Figure 4 are 0.2, 0.1, 0.6, 0.1, which indicates the embedding of current trajectory is more similar to T_1 than T_2 and T_3 . $\mathbf{mix} \in \mathbb{R}^{d \times 1}$ is the weighted sum of the $\mathbf{G_t}$ to concatenate to $\hat{\mathbf{c_t}}$. Finally, a fully connected layer transforms $\mathbf{c_t^{cat}}$ to fit the current state and generate the spatial attention cell state $\mathbf{c_t^{his}}$. Once $\mathbf{c_t^{his}}$ is generated, we can combine it with the intermediate cell state $\hat{\mathbf{c_t}}$ to get the final cell state $\mathbf{c_t}$ by Equation 4.

2) Spatial Memory Writer: The grid cell embeddings in M should be updated during the training process. At each step, the writer directly performs sparse updating of the corresponding entry in the memory M based on s_t:

$$\mathbf{M}(X_a)_{new} = \sigma(\mathbf{s_t}) \cdot \mathbf{c_t} + (1 - \sigma(\mathbf{s_t})) \cdot \mathbf{M}(X_a)_{old}.$$

By this equation, the grid cell embeddings are essentially the weighted average of processed pass-by trajectories. Note that we are using the same spatial gate s_t for both the reader and writer. It is because s_t reflects not only the confidence level that c_{his} is useful for the current input, but also how much information in the current input is suitable for updating $\mathbf{M}(X_g)$. Another benefit of sharing the same gate is that it limits the number of parameters of our model.

V. SEED-GUIDED NEURAL METRIC LEARNING

In this section, We first describe the metric learning procedures of NEUTRAJ and then present the weighted sampling and optimization method to learn the model parameters.

A. Metric Learning Procedures of NEUTRAJ

Figure 1 illustrates our NEUTRAJ model that uses neural networks to embed trajectories into d-dimensional space and approximates the similarity function. As shown, the core of NEUTRAJ is the spatial memory-augmented RNN encoder, which generates latent vector representation for any trajectory. For an input trajectory, the final hidden state of our RNN encoder is used as the trajectory representation.

Given a pool of seed trajectories $\mathfrak S$ and their distance matrix $\mathbf D$, NeuTraj normalizes $\mathbf D$ to a similarity matrix $\mathbf S$ and uses the $\mathbf S$ as guidance. For any two input trajectories T_i and T_j $(i,j\in[1,...,N])$, the RNN encoder is able to project them to two d-dimensional vectors $\mathbf E_i$ and $\mathbf E_j$. Our goal is to learn the network parameters such that similarity between $\mathbf E_i$ and $\mathbf E_j$ is close to the original trajectory similarity $f(T_i,T_j)$. But directly fit all similarities in $\mathbf S$ is intractable and will lead to overfitting. We need select discriminative pairs for optimization. Formally, the loss function of NeuTraj is the weighted sum of MSE of all pairs: $min\sum_{k=1}^K w_k \cdot (f(T_i,T_j)-g(T_i,T_j))^2$, where K is the number of discriminative pairs and w_k is the weight of pair k.

To this end, we design a similarity-preserving ranking objective in NeuTraj. Specifically, for any anchor trajectory T_a , we will sample a set of similar neighbors \mathcal{T}_a^s , as well as a set of dissimilar neighbors \mathcal{T}_a^d from the seed trajectories, to form the similar and dissimilar pairs. With the sampled pairs, we design a weighted ranking objective, which encourages the network to learn a regression function for fitting the similarities to S, as well as preserving the ranking order in both similar and dissimilar pairs. In what follows, we introduce our weighted sampling and optimization procedure.

B. Weighted Sampling and Optimization

Our NEUTRAJ model for metric learning is related to the classic Siamese network [5]. The Siamese network uses random sampling method to generate training pairs, and learns a regression function to fit the target measure. However, such a random sampling strategy implies all pairs have the same weight to the total loss. This assumption does not hold for trajectory metric learning as it ignores the spatial proximity between trajectories. Given one anchor trajectory, we need to focus on the most similar trajectories and the most dissimilar ones, because they are more discriminative than the others. Directly using random sampling and treating all pairs equally can lead to slow convergence and suboptimal accuracies.

To remedy the above problem, we propose a distance-weighted sampling and optimization strategy. We first transform the original distance matrix \mathbf{D} into a normalized similarity matrix \mathbf{S} as follows:

$$\mathbf{S}_{i,j} = \exp(-\alpha \cdot \mathbf{D}_{i,j}) / \sum_{n=1}^{N} \exp(-\alpha \cdot \mathbf{D}_{i,n})$$

where α is a tunable parameter controlling the similarity value distribution. The reason behind the transformation is that the distribution of raw distances often obey to power-law distributions and the magnitude of the distance can span a large range. The transformation is essentially a smoothing operation which brings the similarity values into the range [0,1] and smooths the distribution.

Inspired by [21], our distance-weighted sampling procedure works as follows. We take trajectories in the pool of N seeds as anchor trajectories sequentially. For one anchor trajectory T_a , we take the corresponding row from the similarity matrix ${\bf S}$ as the importance vector ${\bf I_a}$. With the entries in ${\bf I_a}$ as importance weights, we sample n distinct trajectories as similar samples: $\mathscr{T}_a^s = \{T_1^s,...,T_n^s\}$. Conversely, we sample another n dissimilar samples $\mathscr{T}_a^d = \{T_1^d,...,T_n^d\}$ using the entries in ${\bf 1}-{\bf I_a}$ as importance weights. Then we rank the similar samples with the decrease of its similarity to T_a and rank dissimilar samples with the increase order. Finally, we obtain 2n pairs for T_a .

After sampling, we generate the trajectory embeddings and define the pair-wise similarities for the anchor trajectory over the similar and dissimilar pairs as follows:

$$\hat{\mathbf{S}}_{a}^{s} = \hat{\mathbf{S}}(T_{a}, \mathcal{T}_{a}^{s}) = [g(T_{a}, T_{1}^{s}), ...g(T_{a}, T_{n}^{s})]$$

$$\hat{\mathbf{S}}_{a}^{d} = \hat{\mathbf{S}}(T_{a}, \mathcal{T}_{a}^{d}) = [g(T_{a}, T_{1}^{d}), ...g(T_{a}, T_{n}^{d})]$$
(7)

where $g(T_i, T_j) = \exp(-\text{Euclidean}(\mathbf{E_i}, \mathbf{E_j}))$ computes the similarity between two trajectory embeddings, and \mathbf{E} is the embedding of the corresponding trajectory.

Coupled with weighted sampling, we propose a weighted ranking loss which is motivated by list-wise ranking [7] and Mean Reciprocal Rank [22]. Given a ranked list of n sampled trajectories, we set their ranking weights as $\mathbf{r}=(1,1/2,...,1/l,...,1/n)$ and normalize the weights by $\sum_{l=1}^n r_l$. For the n similar pairs, their weights decrease with the ranking order, namely the most similar trajectory in \mathcal{T}_a^s is regarded as the most important. Thus we define the loss for similar pairs of T_a as:

$$L_a^s = \sum_{l=1}^n r_l \cdot (g(T_a, T_l^s) - f(T_a, T_l^s))^2$$
 (8)

where $f(T_i, T_i)$ is the ground truth similarity of (T_i, T_i) .

For dissimilar pairs, it is not reasonable to focus more on fitting the similarity value. Instead, we design a margin loss to separate dissimilar trajectories from the anchor trajectory:

$$L_a^d = \sum_{l=1}^n r_l \cdot [\mathbf{ReLU}(g(T_a, T_l^d) - f(T_a, T_l^d))]^2$$
 (9)

The $\mathbf{ReLU}(x) = max(0,x)$ function defines the margin loss as follows: when $g(T_a, T_l^d) - f(T_a, T_l^d) < 0$, $L_a^d = 0$, meaning that dissimilar sample is faraway enough from the anchor trajectory in the embedding space; when $g(T_a, T_l^d) - f(T_a, T_l^d) > 0$, $L_a^d > 0$, the embeddings should be adjusted to enlarge the embedding-based distance of the dissimilar sample of the anchor. Finally, the loss for the given $\mathfrak S$ is the sum of the similar and dissimilar samples over all N seeds.

$$L_{\mathfrak{S}} = \sum_{a \in [1, \dots, N]} (L_a^s + L_a^d)$$

Since all the modules and the loss functions are differentiable, all the parameters in NEUTRAJ can be learned in an end-to-end manner. In the training process, we update the parameters with back-propagation through time (BPTT) algorithm and employ Adam optimizer for stochastic optimization.

VI. DISCUSSIONS

A. Complexity Analysis

The time complexity of NeuTraj for computing the similarity of a trajectories pair includes two parts: the embedding part and the distance computation part. For embedding, the computation is linear to the number of recurrent operations. In one time step, the higher-order term of complexity in classic recurrent units is $(m+1)*d^2$, where m is the number of gates, e.g., m=3 for LSTM. In SAM units, extra computation cost is involved by a new gate and spatial attention reader which complexity is also quadratic of d. For a pair of trajectories, d is a constant and the complexity of distance computation in the embedding space is a constant. The overall time complexity of NeuTraj is thus linear in the length of the trajectories.

For a trajectory database, the trajectories embeddings only need to be computed once. When new trajectory similarity query is conducted, we generate the embedding of new trajectory and preform search based on the distance of embeddings. So the computation is linear with the size of search space, which makes NEUTRAJ suitable for large dataset.

B. Theoretical Explanation

Similarity measures are used for mapping trajectory to a metric space. In order to make NEUTRAJ general for various of distance measures, we employ a recurrent neural network, which is well known to be a sufficient approximator for an arbitrary mapping function [17], to learn an approximate metric spaces via trajectory embeddings. However, the effects of the neural network module is double-edged. While benefiting the generic property, NEUTRAJ has no provable theoretical guarantee of accuracy. We empirically find that NEUTRAJ can learn a similarity preserving metric space with small size of seed trajectories.

VII. EXPERIMENT

A. Experimental Settings

- 1) Datasets.: Our experiments are based on two public trajectory datasets in two cities: Beijing and Porto. The first dataset [33], referred Geolife, consists of 17,621 trajectories of human mobility from 2007 to 2010. The second dataset [23] consists of over 1.7 millions of taxi trajectories from 2013 to 2014. To moderate the dimensions of M, we choose trajectories in the center area of the city and discretize the area into $50m \times 50m$ grid cells. Then, we remove the trajectories less than 10 records. After such prepossessing, we obtain 8203 trajectories in Geolife and 601,071 trajectories in Porto¹.
- 2) Experimental Protocol: To evaluate the performance of NEUTRAJ, we study top-k similarity search problem on both Geolife and Porto datasets and evaluate NEUTRAJ under four distance measures: the Fréchet distance, the Hausdorff distance, Edit distance with Real Plenty(ERP) and Dynamic Time Warping (DTW). The first three are metric, namely the distance is symmetric and satisfies the triangle inequality. We thus learn the models to approximate the metrics directly. However, the DTW distance is not a metric. Experiments on DTW explore the performance of NEUTRAJ on non-metric similarity measure.

The ground-truth of the problem is the exact top-k results based on the accurate similarity. For Geolife, we compute the accurate similarity of all trajectory pairs and random choose 20% trajectories as the seeds to train NEUTRAJ. Additionally, 10% trajectories are used for tuning parameters and 70% are used for testing. For Proto, due to the enormous trajectories, it is impractical to compute the exact similarity of all trajectory pairs directly. We random choose 10k trajectories to compute the similarity and follow the experimental protocol as the same as Geolife. The performance comparison, efficiency study and parameter sensitivity study of top-k similarity search are shown in VII-B VII-C and VII-D. In addition, based on the well-trained model on 10k Proto trajectories, we conduct a case study of entire Proto dataset(reported in VII-E) to show the efficiency and accuracy of NEUTRAJ on large dataset.

To evaluate the effectiveness of NEUTRAJ for computing pair-wise similarities, we conduct trajectory clustering experiments on both datasets and compare the difference of cluster results between exact similarities and embedding-based similarities. We randomly sample 10k trajectories from Porto, and compute the exact pair-wise similarities as the ground truth. We utilize DBSCAN, which is the most widely used density based clustering algorithm on spatial data, to obtain clustering results and evaluate the difference under four the clustering metrics: Homogeneity, Completeness, V-measure, and Adjusted Random Index. Result of trajectory clustering is shown in VII-F

We also conduct a zero shot learning task to test whether NEUTRAJ works well on a city which has no available trajectories but just the road networks. Based on the road networks

¹Code and data available at https://github.com/yaodi833/NeuTraj

TABLE II PERFORMANCE COMPARISON FOR DIFFERENT METHODS ON FRÉCHET, HAUSDORFF, ERP AND DTW DISTANCES.

Note: HR is the hitting ratio; R10@50 is the top-50 recall for the top-10 ground truth; δ_{H10} is the distortion of average distance on the top 10 results; δ_{R10} is the top 10 recall in top 50 result. The ground truth of top-10 average distance of Fréchet and Hausdorff distances are: 1044m and 730m(Geolife), 1044m distortion of average distance and 730m(Geolife); 935m and 679m(Porto); the element unit of $\delta_{H10}/\delta_{R10}$ is meters.

			F	réchet			Ha	usdorff			ERP			DTW	
Data	Method	HR@10	HR@50	R10@50	$\delta_{\mathrm{H}10}/\delta_{\mathrm{R}10}$	HR@10	HR@50	R10@50	$\delta_{\mathrm{H}10}/\delta_{\mathrm{R}10}$	HR@10	HR@50	R10@50	HR@10	HR@50	R10@50
	AP	0.2374	0.2542	0.5290	213/87	0.2967	0.3180	0.5363	217/113	_	_	_	0.3870	0.4268	0.7139
Geolife	Siamese	0.4631	0.6032	0.8121	162/34	0.3120	0.4236	0.6640	199/69	0.5787	0.7363	0.8964	0.2680	0.4582	0.6172
	NEUTRAJ	0.4947	0.6786	0.8403	84/18	0.3691	0.4870	0.7416	152/42	0.6137	0.7780	0.9424	0.3067	0.4832	0.6513
	AP	0.2542	0.2851	0.5520	208/79	0.2832	0.2966	0.5620	201/86	_	_	_	0.3798	0.4160	0.7010
Porto	Siamese	0.4740	0.5802	0.7970	128/27	0.3834	0.4999	0.7760	165/48	0.4982	0.6893	0.9043	0.3832	0.4804	0.7602
	NEUTRAJ	0.5225	0.6351	0.8292	89/ 8	0.4372	0.5714	0.8089	101/15	0.5427	0.7297	0.9277	0.4370	0.5613	0.8396

 ${\it TABLE~III} \\ {\it Results~of~ablation~experiments~for~different~methods~on~Fr\'echet,~Hausdorff,~ERP~and~DTW~distances.} \\$

			F	réchet			Ha	usdorff			ERP			DTW	
Data	Method	HR@10	HR@50	R10@50	$\delta_{\mathrm{H}10}/\delta_{\mathrm{R}10}$	HR@10	HR@50	R10@50	$\delta_{\mathrm{H}10}/\delta_{\mathrm{R}10}$	HR@10	HR@50	R10@50	HR@10	HR@50	R10@50
	NT-No-WS	0.4736	0.6353	0.7996	139/27	0.3338	0.4393	0.6273	169/55	0.5880	0.7170	0.8686	0.2591	0.4610	0.6260
Geolife	NT-No-SAM	0.4842	0.6483	0.8198	117/23	0.3574	0.4607	0.7219	157/46	0.6090	0.7537	0.9291	0.2881	0.4792	0.6482
	NEUTRAJ	0.4947	0.6786	0.8403	84/18	0.3691	0.4870	0.7416	152/42	0.6137	0.7780	0.9424	0.3067	0.4832	0.6513
	NT-No-WS	0.4990	0.5883	0.7981	102/10	0.4190	0.5628	0.7909	140/33	0.5192	0.6920	0.8917	0.3930	0.5013	0.7919
Porto	NT-No-SAM	0.5154	0.6121	0.8171	92/10	0.4238	0.5691	0.8033	126/16	0.5382	0.7111	0.9107	0.4238	0.5425	0.8148
	NEUTRAJ	0.5225	0.6351	0.8292	89/ 8	0.4372	0.5714	0.8089	101/15	0.5427	0.7297	0.9277	0.4370	0.5613	0.8396

in Beijing [32], we simulate 6000 synthetic trajectories as the seeds for training and test NEUTRAJ with the real trajectories in Geolife. Result of zero short learning is presented in VII-G.

- 3) Compared Methods: For the studied four measures, we compare NEUTRAJ with four baselines, which can be roughly divided into three categories:
- Approximate algorithms: Except ERP which has no approximate algorithm, each of the three measures has several approximate algorithms to fast compute them. We compare with the state-of-the-art approximate algorithms from [12] which is used for computing both Fréchet and DTW distances, and [4] which is used for Hausdorff distance. We call these algorithms as AP in general for all the distance measures.
- Siamese Network [24]: This category is a metric learning approach based on the Siamese network. We instantiate the Siamese network with LSTM backbone, and denote it as Siamese.
- Ablations: Finally, we include two kinds of ablations of NEUTRAJ. (1) The weight sampling in NEUTRAJ is replaced by random sampling to test the effectiveness of distance-weighted ranking loss. We denote this variant as NT-No-WS, respectively. (2) We replace the SAM unit with standard LSTM, denoted as NT-No-SAM, to test the effects of the proposed spatial attention memory mechanism.
- 4) Evaluation Metrics: We use three different metrics for performance evaluation. The first is the top-k hitting ratio, which examines the overlap percentage of the top-k results and the ground truth. We report the hitting ratio for both top-10 (HR@10) and top-50 searches (HR@50). The second is the top-50 recall for the top-10 ground truth (R10@50). This one evaluates how many of top 10 ground-truth trajectories are recovered by the top 50 lists produced by different methods. The third metric is the distortion of average distance for the top-10 results, denoted as δ_{H10} and δ_{R10} . δ_{H10} is computed based on the top-10 trajectories from the results and δ_{R10}

is based on the top-10 most similar trajectories from top-50 results. They measure the distortion of average exact distances between the query trajectory and the top-10 search results. The smaller the distance is, the stronger a method performs.

5) Parameter Settings: The key parameters in NEUTRAJ include: (1) the embedding dimension d; (2) the scan width w of the attention memory reader. We have tuned d by the grid search in range $\{16,32,64,128,256\}$. In general, the performance increases with d and gradually stabilizes when it is large enough. For w, we tuned it in $\{0,1,2,3,4\}$ and found that it had an optimal value as w=2 on both datasets. Finally, we set d=128 and w=2. In addition, we set the batch sizes as 20 and the sampling size n as 10. For the compared methods, we tuned their parameters to obtain the best performance in our datasets. We will also report the parameter study results in Section VII-D.

B. Performance Comparison

Table II shows the performance of different methods for the top-k similarity search task. As shown, on both datasets, NEUTRAJ significantly outperforms all the baseline methods in most of metrics. Take the Fréchet distance as an example. Compared with state-of-the-art approximate algorithms (AP), the variant of NEUTRAJ, i.e., NEUTRAJ achieve more than two time higher hitting ratios, about 70% gain in R10@50, and about 69% reductions in average distance. Such huge improvements is impressive given the fact that NEUTRAJ does not rely on any hand-crafted heuristics but learns the similarity function automatically from seed trajectories. The superiority of NEUTRAJ over the Siamese network is also obvious in all the four metrics. While both methods employ neural metric learning for approximating the similarity function, NEUTRAJ has two advantages over the Siamese network. First, the weighted sampling and ranking loss can yield more distinguishing trajectories and training loss compared to Siamese network. Second, the spatial attention memory module models the correlation between spatially close trajectories, which is

very beneficial for generating trajectory embedding of high quality. Among the four distances, the performance of DTW is inferior to other three. The reason is that the distance of learnt embeddings is a metric while DTW is not. This systematic error limits NEUTRAJ effectively performing on DTW.

The results of ablation experiments are shown in Table III. Comparing NeuTraj with its ablations, one can further see the effectiveness of the two major modules in NeuTraj. Continue using the Fréchet distance on the Geolife dataset as an example. We observe: (1) by including the SAM module, NeuTraj improves HR@10 of NT-No-WS from 0.46 to 0.47; and (2) by including the weighted sampling and optimization module, NeuTraj improves the HR@10 of NT-No-WS from 0.47 to 0.49. The trend is similar for the Porto dataset and other three similarity measures.

The absolute values of HR@10 and HR@50 seem not very high on both datasets. The reason is that trajectories in both datasets have lots of near-duplicate instances. This can be observed from δ_{H10} , which measures the spatial closeness for both the top-10 ground truth and the generated top-10 list.

C. Efficiency Study

In this subsection, we study the efficiency of NEUTRAJ. We first report its time cost for online similarity search, and then report the offline time cost for training the NEUTRAJ model. The experiments are conducted on a machine with Inter Xeon E5 @2.20GHz CPU and one Nvidia P100 GPU.

 $\label{total cost for online similarity search without index.}$ Time cost for online similarity search without index.

Method	1k	5k	10k	200k
		Fı	réchet	
BruteForce	8.712s	41.876s	84.480s	1639.834s
AP	1.840s	11.319s	23.107s	532.652s
NT-No-SAM	0.461s	0.471s	0.489s	1.576s
NEUTRAJ	0.461s	0.470s	0.490s	1.574s
		Ha	usdorff	
BruteForce	0.238s	1.416s	2.981s	51.642s
AP	0.127s	0.154s	0.179s	3.426s
NT-No-SAM	0.026s	0.046	0.072s	1.133s
NEUTRAJ	0.024s	0.047	0.073s	1.131s
]	ERP	
BruteForce	0.409s	1.982s	3.807s	73.054s
NT-No-SAM	0.027s	0.046s	0.081s	1.154s
NEUTRAJ	0.026s	0.047s	0.081s	1.152s
		Ι	DTW	
BruteForce	0.305s	1.482s	3.070s	59.054s
AP	0.119s	0.142s	0.185s	4.021s
NT-No-SAM	0.023s	0.044s	0.066s	1.028s
NEUTRAJ	0.021s	0.043s	0.067s	1.027s

1) Time Cost of Online Similarity Search: Similarity Search without Index. Table IV shows the time cost of NEUTRAJ for performing top-k similarity search with different sizes. Specifically, from the test set of Porto, we randomly sample four sub-corpora with sizes 1K, 5K, 10K, and 200K respectively. Then we use NEUTRAJ to perform top-50 similarity for each trajectory and re-rank the 50 trajectories by calculating their accurate distance. Table IV reports the average time cost for processing one query. We compare it with the BruteForce method that directly computes the exact distances

following the definition, state-of-the-art approximate algorithms (AP), as well as the neural network based methods(NT-NO-SAM). We omit the results of NT-NO-WS and Siamese methods because the time cost of NT-NO-WS and Siamese methods are analogous to NEUTRAJ and NT-NO-SAM in online search procedure. As shown in Table IV, for all the four measures, NEUTRAJ achieves 50x-1500x speedup over BruteForce and 2x -500x speedup over existing approximate algorithms. The speedup ratios are especially significant for large datasets. The time cost of ablation methods NT-NO-SAM is very close to NEUTRAJ because the embedding time difference of one search trajectory is small.

 $\label{thm:table V} TIME\ COST\ FOR\ ONLINE\ SIMILARITY\ SEARCH\ WITH\ INDEX$

Method	1k	5k	10k	200k
	В	ounding B	ox R-tree	Index
BruteForce	5.526s	27.802s	54.558s	1070.433s
AP	0.438s	1.731	4.372s	62.853s
NEUTRAJ	0.005s	0.029	0.056s	0.868s
# of involved trajectories	675	3377	6736	134051
	(Frid-based	Inverted I	ndex
BruteForce	5.633s	27.793s	54.993s	1098.042s
AP	0.460s	1.911	4.722s	66.072s
NeuTraj	0.006s	0.030	0.065s	1.173s
# of involved trajectories	685	3424	6834	136201

Similarity Search with Index. In this experiment, we employ two widely used indexing techniques: 1) bounding box r-tree; 2) grid based inverted index, and random select 200 query trajectories to examine the elastic character of NEUTRAJ. Under the Fréchet distance, we compare indexextended NEUTRAJ with two baselines: brute force method and approximate algorithm, on various sub-corpora. The average time costs are reported in Table V. We find that NEUTRAJ outperforms the baselines under the two indexing structures and achieves over 30x speedup compared with the approximate algorithm. Similar trend can be observed on other measures.

 $\label{eq:table_vi} \mbox{TABLE VI} \\ \mbox{Time cost for offline model training.}$

Methods		Model Trair	1	Embedding					
Methods			t_{total}	200k					
Siamese	164s	71	11644s	411s					
NEUTRAJ	285s	15	5130s	639s					
	Ablations Experiments								
NT-No-SAM	168s	15	2520s	412s					
NT-No-WS	283s	20	5660s	636s					

2) Time Cost of Offline Training and Embedding: Offline Training Time. Table VI shows the comparison of offline training time on the Porto dataset under the Fréchet distance for other similarity measures. For 2,000 training trajectories, NEUTRAJ converges within 20 epochs. The training time of one epoch is around 5 min for NEUTRAJ, and thus the entire training time of NEUTRAJ is less then 2 hours. In contrast, the Siamese network takes more than 60 epochs to converge, which is about 3X slower than NEUTRAJ. We also compare the convergence rate of NEUTRAJ and NT-No-SAM in Figure 5 and observe that NEUTRAJ has higher convergence rate than NT-No-SAM because SAM module captures useful information from processed trajectories.

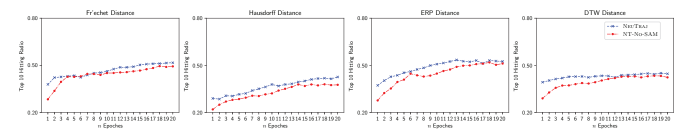


Fig. 5. The convergence curve of NEUTRAJ and NT-NO-SAM on Fréchet, Hausdorff, ERP and DTW with respect to 20 epochs.

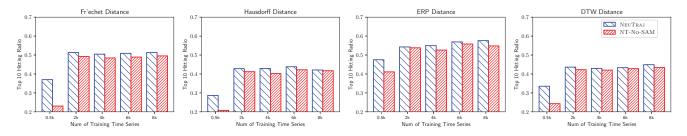


Fig. 6. HR@10 of NEUTRAJ and NT-NO-SAM on Fréchet, Hausdorff and DTW with varying training data size.

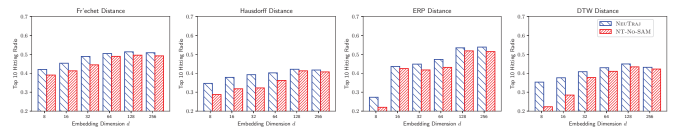


Fig. 7. HR@10 of NEUTRAJ and NT-No-SAM on Fréchet, Hausdorff and DTW with varying embedding size d.

Offline Embedding Time. Another part of offline training is the embedding procedure which generates the embeddings of trajectories by using the well-trained model. In this experiment, we set the embedding batch size as 2000 and report the time cost of embedding 200k trajectories. The results of all neutral network based methods are shown in Table VI. We can easily find that SAM unit based methods(NEUTRAJ, NT-NO-WS) are little slower than standard unit based method(NT-NO-SAM, Siamese). The reason is SAM unit need more calculation for finding useful information in memory tensor.

D. Parameter Sensitivity Study

In this subsection, we evaluate the sensitivity of NEUTRAJ on three parameters: the training data size, the scan width w, and the embedding dimension d. Due to the space limitation, we only shows the experimental result on the Porto dataset.

1) The sensitivity of training data size: We first investigate the effect of the number of seed trajectories on the performance of NEUTRAJ. Figure 6 shows the results of NEUTRAJ and its ablation NT-NO-SAM for the four measures as we vary the training data size from 500 to 8,000 on Porto. As shown, the performance of NEUTRAJ becomes relatively stable when there are more than 2,000 training trajectories. We

also observe that SAM-based models outperform the NT-NO-SAM models. Another interesting fact is that NEUTRAJ is more robust than NT-NO-SAM with sparse training data. As shown, when the number of training trajectories is only 500, the performance gap between NEUTRAJ and NT-NO-SAM is particularly large. This is because SAM employs a memory tensor to memorize useful information from processed trajectories.

- 2) The sensitivity of embedding dimension d: We proceed to study the effect of the embedding dimension d on the performance of NeuTraj. Figure 7 illustrates HR@10 as we vary d from 8 to 256. As shown, the performance of NeuTraj and its variants first increases and then drops slightly. The reason is the parameter d controls the complexity of NeuTraj. When d increases, the model enjoys more expressive power to capture the intrinsic structures of trajectories. However, when d is too large, the model suffers from over-fitting due to the limited size of training data.
- 3) The sensitivity of scan width w: Finally, the scan width w in the SAM module is a key parameter that controls the exploration spread for historical trajectories. As shown in Figure 8, with the increase of w, the HR@10 for all methods first increases and then slightly drops. This phenomenon is

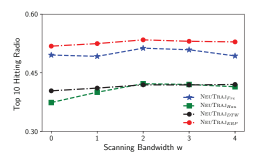


Fig. 8. HR@10 of NEUTRAJ with varying w.

attributed to two reasons: (1) as w increases, more information tends to be accessed by the SAM reader, which is useful for encoding the current trajectory in the initial stage; (2) when w is too large, the information of some non-relevant trajectories will be inevitably incorporated. Even if the attention mechanism in NEUTRAJ can help reduce this effect, the performance of the model can still be harmed.

E. Case Studies

We use the entire Porto dataset to perform several case studies to intuitively examine the top-k search results of our model. For this purpose, we randomly choose several trajectories and retrieve their top-5 neighbors under the Fréchet distance. Due to the space limitation, Table VII only shows the results of two representative trajectories: T_{91} and T_{65} . For each query, we plot both the top-3 ground truth trajectories as well as the top-3 trajectories retrieved by NEUTRAJ.

The result shows that NEUTRAJ is quite effective on both short(T_{91}) and long(T_{65}) trajectory: the results returned by NEUTRAJ match the ground truth very well, and the distortions of top-5 average distance δ_{H5} are very small. Moreover, by training with the weighted ranking loss, NEUTRAJ preserves the ranking order of trajectories.

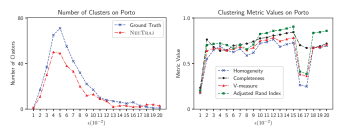


Fig. 9. Trajectory Clustering Result.

F. Result of Trajectory Clustering

In this experiment, we aim at exploring the effectiveness of NeuTraj via trajectory clustering. Due to the space limitation, we only show the clustering result of DBSCAN using Fréchet distance on Porto dataset and compare the two cluster results which are generated by the ground truth distance and embedding based distance. As shown in Figure 9, fixing the minimum points as 10, the number of clusters of the two results change similarly with the increase of ϵ . Best values

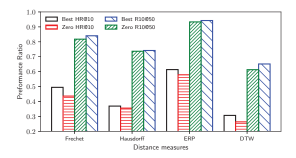


Fig. 10. Illustration of zero-shot learning results on Geolife dataset. Best is the best performance NeuTraj can achieve on real dataset and Zero is the performance trained with synthetic data.

of the evaluation metrics are more than 0.8, which indicates NEUTRAJ works well on this task. Similar trends are observed on other three distance measures and Geolife dataset.

G. Evaluation on Zero-Shot Learning

In the final set of experiments, we are interested in applying NEUTRAJ for zero-shot learning scenarios. Specifically, the NEUTRAJ model relies on a real trajectory database and sampling seeds from the database as guidance. It is interesting to investigate: how NEUTRAJ perform if there are no real seed trajectories? For this purpose, we extend NEUTRAJ for the zero-shot learning scenario. We assume there exists no trajectory databases but only a road network of the target area. Then we generate a bunch of simulated trajectories as our seeds and train the NEUTRAJ model for computing the similarity for a pair of real trajectories. Based on the road network in Beijing [32], we generate 6,000 synthetic trajectories by employing random walk on road node graph and interpolating coordinates between the nodes. Then we take the synthetic trajectories as the seeds to train NEUTRAJ, and test it with the real trajectories from Geolife. Figure 10 shows the HR@10 and R10@50 of NEUTRAJ.

Impressively, even using synthetic trajectories and their distances as guidance, NEUTRAJ can still achieve around 0.7 recall for all the four metrics. Such a phenomenon indicates that NEUTRAJ can be applied to scenarios even when no real trajectory databases are available.

VIII. CONCLUSION

We proposed a seed-guided neural metric learning approach NEUTRAJ that is fast, accurate, generic and elastic for trajectory similarity computation. Its novelty lies on two aspects: (1) a spatial attention memory (SAM) module that can augment existing RNN architectures to capture the correlations between trajectories; and (2) a distance-weighted ranking loss that effectively leverages seed information to learn trajectory embeddings of high quality. Experiments on two real-life datasets have shown that NEUTRAJ can effectively accelerate on various distance measures, while producing more accurate function approximations over state-of-the-art baselines.

Several interesting problems exist for future exploration. First, NEUTRAJ mainly focuses on two-dimensional trajectories. It is interesting to extend NEUTRAJ for trajectories

Result of T_9	1: HR@10: 0.7;	HR@50: 0.78;	H10@R50: 0.9;	$\delta_{H5} = 4\text{m}; \delta$	t_{H10} = 4m; δ_{R1}	$_{0} = 2m$
Query Trajectory		Top-3 ground truth			Top-3 of NEUTRA.	l.
Trai 91	Rank 1 in GT	Rank 2 in GT	Rank 3 in GT	*Rank 3 in GT Rank 1 in Neutraj	Rank 6 in GT Rank 2 in Neutraj	e Bank 7 in GT Rank 3 in NeuTraj Traj 44081
Result of T_{65} :	HR@10: 0.4: H	R@50: 0.52; H	10@R50: 0.8; δ	$H_5 = 296 \text{m}; \delta$	$H_{10} = 236 \text{m}; \delta_{I}$	$R_{10} = 62m$
	,	11000, 0.02,	10 € 1650. 0.0,	$H_0 = 2$	$H_{10} = 250 \text{m}, 0_I$	210 - 02111
Query Trajectory		Top-3 ground truth		H 5 - 250III, 0	Top-3 of NEUTRAL	

with time dimension. Second, NEUTRAJ is designed for accelerating similarity computation of trajectory pairs and only suits for similarity based queries. It would be interesting to investigate new models to handle other types of query, such as range query and boolean query.

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