A Scalable Framework for Trajectory Prediction

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Abstract—Trajectory prediction (TP) is of great importance for a wide range of location-based applications in intelligent transport systems such as location-based advertising, route planning, traffic management, and early warning systems. In the last few years, the widespread use of GPS navigation systems and wireless communication technology enabled vehicles has resulted in huge volumes of trajectory data. The task of utilizing this data employing spatio-temporal techniques for trajectory prediction in an efficient and accurate manner is an ongoing research problem. Existing TP approaches are limited to short-term predictions. Moreover, they cannot handle a large volume of trajectory data for long-term prediction. To address these limitations, we propose a scalable clustering and Markov chain based hybrid framework, called Traj-clusVAT-based TP, for both short-term and long-term trajectory prediction, which can handle a large number of overlapping trajectories in a dense road network. Traj-clusVAT can also determine the number of clusters, which represent different movement behaviours in input trajectory data. In our experiments, we compare our proposed approach with a mixed Markov model (MMM)-based scheme, and a trajectory clustering, NETSCAN-based TP method for both short- and long-term trajectory predictions. We performed our experiments on two real, vehicle trajectory datasets, including a large-scale trajectory dataset consisting of 3.28 million trajectories obtained from 15,061 taxis in Singapore over a period of one month. Experimental results on two real trajectory datasets show that our proposed approach outperforms the existing approaches in terms of both short- and long-term prediction performances, based on prediction accuracy and distance error (in km).

Index Terms—Large-scale trajectory data, next location prediction, Long-term trajectory prediction, scalable clustering.

I. INTRODUCTION

With the widespread use of Global Positioning System (GPS) devices, smart-phones, Internet of Things [1], and wireless communication technology, it is possible to track all kinds of moving objects all over the world. The increasing prevalence of location-acquisition technologies has resulted in large volumes of spatio-temporal data, especially in the form of trajectories. These data often contain a great deal of information [2], which give rise to many location-based services (LBSs) and applications such as vehicle navigation, traffic management, and location-based recommendations. One key operation in such applications is the route prediction of moving objects.

Vehicle route prediction allows certain services to improve their quality e.g. if the route of vehicles is known in advance, intelligent transportation systems (ITSs) can provide route-specific traffic information to drivers such as forecasting traffic conditions and routing the driver so as to avoid traffic jams. Route prediction also enables location-based advertising, which can advertise certain products/services and special offers to the target commuters most likely to pass through business outlets and stores based on their travel trajectory.

With the advancement in artificial intelligence, various machine learning techniques have now been used for trajectory analysis. Recently, several studies have been carried out on trajectory prediction, particularly after Song et al. [3] demonstrated a 93% potential for predictability in user mobility, which supplied the theoretical basis for location prediction methods. These methods mainly focus on two kinds of prediction models. The first type is the short-term trajectory prediction model, which aims to predict the next-location or a few locations in the near future. These models usually rely on current location and one or two previous locations of an object to predict its next location. The second type is the long-term trajectory prediction model which focuses on location prediction at a more distant future time or on complete route prediction. These models generally rely on an available partial trajectory of a moving object to predict the complete trajectory.

In urban areas, vehicle trajectories are usually constrained to a complex road network with many parallel and perpendicular road segments and intersections, which makes their time progression very irregular. Due to the uncertainty of moving objects, most of the existing trajectory prediction methods only focus on predicting short-term partial trajectories. They have poor prediction accuracy for long-term trajectory predictions and they do not work well for estimating continuous and complete trajectories.

The sheer amount of vehicle trajectory data, if analyzed effectively, can significantly improve route prediction performance. However, it is challenging to carry out trajectory prediction from a large amount of trajectory data. The huge volumes of data to be processed precludes using machine learning based traditional prediction (TP) methods. Most of the existing trajectory prediction schemes cannot handle a large number of trajectories, especially when they span a large area of a road network. Existing TP methods are hybrid in nature and usually employ classical frequent sequential pattern based algorithms, Markov model-based algorithms, or clustering based algorithms to a small amount of training data. Apriori-like sequential pattern methods bear significant computational overhead in mining frequent patterns because they
produce a huge number of candidate sequences and require multiple scans of the database. Markov model approaches such as higher-order Markov models incur significant overhead in terms of computation time and space for storing and learning the mobility model. Similarly, parameter learning algorithms in Hidden Markov models impose significant computational burdens for large-scale trajectory datasets. Existing trajectory clustering based location prediction approaches are not scalable due to distance matrix computations for big data and are not able to handle a large number of overlapping trajectories efficiently. Therefore, most TP methods demonstrated in the literature use synthetic or small to medium size real trajectory datasets. To the best of our knowledge, the largest real dataset was used in [4], consisting of 370 million GPS points. These authors utilized a parallel processing model, MapReduce, in their implementation to handle large datasets.

To address these challenges and overcome the drawbacks of existing TP methods, this paper presents a novel, scalable framework for both short-term and long-term TP, which is suitable for large numbers of overlapping trajectories in a dense road network, typical for major cities around the world. First, we cluster the large trajectory data using a modified version of our two-stage clusiVAT (clustering using improved Visual Assessment of Tendency) algorithm [5], [6], which we call Traj-clusiVAT, implemented for trajectory prediction task. The Traj-clusiVAT algorithm first extracts a smart sample using the Maximin-Random sampling (MMRS) scheme [7], which provides a good representation of the input cluster structure (present in the original data). Then, it uses the improved visual assessment of cluster tendency (IVAT) algorithm to visually determine the number of clusters \( k \) in input data, and subsequently, it partitions the trajectory sample into \( k \) clusters which contain different frequent movement patterns in the trajectory data. Then, the remaining non-sampled trajectories are assigned to one of \( k \) clusters using the nearest prototype rule (NPR). Finally, Markov chain models are constructed from the trajectories in each cluster. These models quantify the movement patterns within clusters, and subsequently, can be used for TP.

Our main contributions in this paper are as follows:

- We propose a novel, hybrid framework for short-term and long-term trajectory prediction, based on a scalable clustering algorithm and Markov chain model, which can utilize a large number of trajectories in a dense road network.
- We implement a modified version of the two-stage clusiVAT algorithm, called as Traj-clusiVAT, to cluster large numbers of trajectories in an accurate and efficient manner. In this regard,
  - We develop a new algorithm to compute a representative trajectory (RT) for each cluster, and subsequently, use it to assign non-sampled trajectories to one of the \( k \) clusters in the nearest prototyping phase.
  - We develop an improved algorithm for nearest prototyping, which assigns each non-sampled trajectory to one of the \( k \) clusters, based on pattern matching and the distance of non-sampled trajectories to the RT of each cluster.
- We compare our proposed algorithm with two TP algorithms: a mixed Markov model (MMM) [8] and a trajectory clustering model NETSCAN [9] based algorithm. Our experiments used two real-life, taxi trajectory datasets including a large-scale taxi trajectory dataset consisting of 370 million GPS traces and 3.28 million passenger trips from 15,061 taxis during the period of one month in Singapore. To the best of our knowledge, this is the first time TP has been performed on such a large number of real-life road network trajectories.

Here is an outline of the rest of this article. Section II discusses related work and limitations. Section III formally defines the problem and presents preliminaries on techniques used in our work. Our novel, hybrid framework, based on Traj-clusiVAT and Markov models, is presented in Section IV and its time complexity is discussed in Section V. Section VI presents our numerical experiments, including computation protocols, datasets and results, followed by conclusions in Section VII.

II. RELATED WORK

Several studies address the problem of trajectory predictions, which includes the problem of short-term prediction such as predicting the next location, and long-term prediction such as future locations or complete route prediction. These methods mainly focus on discovering frequent patterns using various data mining methods. Many of these methods are hybrid in nature and can be broadly classified into three categories: (i) Rule-based learning based approaches (ii) Markov model-based approaches (iii) Clustering-based approaches.

A. Rule-based learning based approaches

Several rule-based methods have been used for location prediction. Morzy [10] implemented a modified version of the PrefixSpan algorithm to extract association rules from a moving object database, and used frequent pattern tree with a matching function to select the best association rule from the database of movement rules. Jeung et al. [11] proposed a hybrid prediction approach, which combines association rules in the form of trajectory patterns with the motion functions of an object’s recent movements, to estimate future locations. Given an object’s recent movement and predictive queries, the best association rule is chosen for prediction. The query processing approaches presented in [11] can only support near and distant-time predictive queries, making them unsuitable for long-term trajectory prediction. Moreover, with the huge number of trajectories, the number of association rules is also huge, which makes association-rule based algorithms impractical for large-scale mobility data.

Monreale et al. [12] built a decision tree that they called a T-pattern Tree, based on the frequent movement patterns extracted using a Trajectory Pattern algorithm, and predicted the next location of a new trajectory based on the best matching functions. However, mining of frequent trajectory patterns is computationally expensive. The method in [12] is similar to the use of association rules as predictive rules in rule-based classifiers. Therefore, this method [12] may result in
a large number of predictive rules for voluminous trajectories. Qiao et al. [13] proposed a TP algorithm, called PrefixTP, which examines only the prefix subsequences, and projects their corresponding postfix subsequences into projected sets. Then, for a partial trajectory, it recursively finds a postfix sequence based on the minimum support count requirement and then declares the most frequent sequential pattern as the most probable trajectory. Finding subsets of trajectory sequential patterns is a recursive mining process, which is also computationally extensive.

B. Markov model-based approaches

Markov models (MMs) have been widely used to mine frequent patterns for route prediction problems. Ishikawa et al. [14] proposed a model to extract mobility statistics, called the Markov transition probability, which is based on a cell-based organization of target space and a Markov chain model, and employed R-tree spatial indices to compute Markov transition probabilities. Simmon et al. [15] presented a Hidden Markov Model (HMM) based probabilistic approach to predict a driver’s intended route and destination through observations of the driver’s habits. Asahara et al. [8] suggested that standard MM and HMM are not generic enough to encompass all types of movement behaviour. They proposed a variant of Markov model, called the mixed Markov-chain model (M3M), as an intermediate model between individual and generic models, for pedestrian movement prediction.

Gambs et al. [16] extended a previously proposed mobility model, named v-Mobility Markov Chain (v-MMC), to incorporate the v previous visited locations. They showed that prediction accuracy increases with v, but increasing v beyond two does not compensate for the significant overhead in terms of computation and space for learning and storing the mobility model. They only considered the sequence of the significant locations, instead of all locations, to build higher order MM. Zhang et al. [17] proposed a group-level mobility modeling method, Gmove, which alternates between two intertwined tasks, user grouping and mobility modeling, and generates an ensemble of HMMs to characterize group-level movement.

Most of the MMs do not consider the discontinuous chain of the hidden states, and therefore, the state retention problem can drastically degrade the accuracy of location prediction system [13]. For the irregular trajectory data, the movement rules cannot be easily represented by Markov models, which may cause loss of continuous location information [13]. Moreover, the HMM approaches use the Baum-Welch algorithm for parameter learning and the Viterbi algorithm to find the most likely sequences of hidden states. These algorithms impose a significant computation burden for large-scale trajectory datasets.

Recently, deep learning techniques such as inverse reinforcement learning (IRL), long short-term memory (LSTM), and recurrent neural networks (RNNs) have been used for modeling vehicle trajectories [18–20]. However, some of these studies are still based on first-order Markov assumption to model the routing decisions for heterogeneous destinations, or they are too shallow which makes the modeling pattern varieties suffer from too few parameters.

C. Clustering based approaches

Some researchers have proposed trajectory clustering based route prediction methods, which partition the trajectories into several clusters representing different motion patterns based on the trajectory similarity. Various clustering approaches [21] using different methods and distance measures between trajectories have been proposed in the literature. Road network constrained trajectory clustering approaches can be classified into two broad categories. The first type uses the traditional clustering approaches such k-means and DBSCAN with specially designed distance measures [22]–[25] for trajectories. The second category of algorithms [9], [26] cluster road segment vehicle frequencies based on density and flow.

Ashbrook et al. [27] presented a system that automatically detected the significant locations from GPS data using k-means clustering, and then incorporated these locations into an MM to predict the next location. Mathew et al. [28] presented a hybrid method for human mobility prediction, which first clusters location histories according to their characteristics, and then trains an HMM for each cluster. A poor prediction accuracy of 13.85% was obtained on a real, large-scale trajectory dataset using this method. Chen et al. [29] proposed a next-location prediction approach combining two clustering models, which cluster the objects based on the spatial locations and trajectories using a similarity metric, respectively, and trains a series of MMs with trajectories in each cluster.

Ying et al. [30] proposed an approach for predicting the next location based on geographic and semantic features of user trajectories. This method requires the calculation of a semantic score for each candidate path, which generally incurs additional overhead when compared with other methods. A probabilistic TP model was proposed in [31] based on two mixture models, a Gaussian Mixture Model (GMM) and a Variational Gaussian Mixture Model (VGMM), optimized using the Expectation Maximization (EM) algorithm. Their method requires the prior selection of the number of Gaussian components and other distribution parameters. They evaluated their method on a small dataset, which consists of only 69 trajectories. Qiujian et al. [4] proposed a spatio-temporal prediction and a next-place prediction model based on an entropy-based clustering approach and HMMs.

Traditional clustering [22]–[25] based prediction methods are not scalable for a large number of trajectories as distance matrix computation is time and space prohibitive. Most of them require the number of clusters to be known in advance, but in practice, it is often unknown, making it difficult for the user to choose the optimal number of clusters for location prediction. Furthermore, the clusters are determined by fixed rules. Some of the road network based clustering approaches [9], [26], though scalable, produce clusters having high intra-cluster variance, which span a large area of a road network.

Most of the work done in the area of trajectory prediction either use synthetic datasets [8], [10], [11], [32] or real datasets with small to medium numbers of data points [12], [29], [33]. Most of them cannot handle big trajectory datasets. There have been several attempts to demonstrate trajectory prediction on real data having a large number of samples. For
example, \[13\] uses a real dataset consisting of 4.9 million trajectories (790 million GPS points) as a population, but only small subsets having a maximum 30,000 trajectories are used in their experiments. To the best of our knowledge, the largest real dataset used was in \[15\], consisting of 37 million GPS points. They utilized \[2\] the MapReduce model in their implementation to handle large datasets. In this paper, we use the GPS traces of 15,061 taxis in Singapore over a period of one month. We extract 3.28 million passenger trajectories consisting of 370 million GPS logs from this data for trajectory prediction task. To the best of our knowledge, this is the first time trajectory prediction task has been performed on such a large number of real-world road network trajectories.

### III. Preliminaries

In this section, we introduce some basic terms and definitions, which are required in the sequel.

#### A. Road Network and Trajectories

We represent the road network as an undirected graph

\[ G_{RN} = (V, E), \]

comprising a set \( V \) of intersections or nodes of the road network with a set \( E \) of road segments or edges, \( R_i \in E \) such that \( R_i = (r_{i_a}, r_{i_b}) \), where \( r_{i_a}, r_{i_b} \in V \) and there exists a road between \( r_{i_a} \) and \( r_{i_b} \). The edge \( R_i \) is given a weight equal to the length of \( R_i \). For such a road network, we define the following:

**Definition 1.** (Trajectory): A trajectory \( T \) of length \( l \) is a time ordered sequence of road segments (RS), \( T = (R_1, R_2, \ldots, R_l) \), where \( R_j \in E, 1 \leq j \leq l \), and \( R_j \) and \( R_{j+1} \) are connected.

**Definition 2.** (Sub-Trajectory): A trajectory \( T = (R_1, R_2, \ldots, R_p) \) is a sub-trajectory of sequence \( T = (R_1, R_2, \ldots, R_l) \), \( p \leq l \), if there are integers \( i_1, i_2, \ldots, i_p \) \( 1 \leq i_1 < i_2 < \ldots < i_p \), \( j_1, j_2, \ldots, j_p \) \( 1 \leq j_1 < j_2 < \ldots < j_p \), \( |j_1, j_2, \ldots, j_p| = |i_1, i_2, \ldots, i_p| = |j_1, j_2, \ldots, j_p| = |i_1, i_2, \ldots, i_p| = |j_1, j_2, \ldots, j_p| \). Then \( T^* \) is called a sub-trajectory of \( T \), denoted by \( T^* \subseteq T \).

**Definition 3.** (Frequent Road Segment): A Frequent road segment (FRS), \( R_{FRS} \), in a trajectory set is a segment that contains at least \( MinT \) percentage of trajectories of the set passing through the segment, otherwise, the segment is labeled as "non-FRS". The percentage \( MinT \) is a tunable parameter and we call it the FRS threshold.

**Definition 4.** (Partial Trajectory): A partial trajectory \( T^p \) is a sub-trajectory of a given trajectory \( T \) if and only if their sequences start from the same segment.

**Definition 5.** (Source Segment): The segment from which a trajectory \( T \) originates is called the Source Segment (SS), \( R_{SS} \), and the start node of \( T \) is called the Source Node (SN) of that trajectory. For a trajectory \( T = (R_1, R_2, \ldots, R_l) \), the road segment \( R_1 \) is \( R_{SS} \). Node \( r_{i_a} \) is the SN, if \( R_2 \) has node \( r_{i_b} \), else \( r_{i_b} \) is SN, where \( R_1 = (r_{i_a}, r_{i_b}) \), and \( r_{i_a}, r_{i_b} \in V \).

**Definition 6.** (Frequent Source Segment): The SS which is FRS, is called Frequent source segment (FSS), \( R_{FSS} \).

**Definition 7.** (Problem Definition): Assume that a historical trajectory database, containing \( N \) trajectories, denoted by \( T = \{T_1, T_2, \ldots, T_N\} \) is given. Then, for a given partial trajectory \( T^p = (L_1, L_2, \ldots, L_m) \), the goal is to predict the future road segments \( L_0 \), where \( i, m \in Z \) and \( i \geq m + 1 \).

#### B. Distance Measure (trajDTW)

Most of the existing distance measures for trajectory similarity are not suitable for a large number of overlapping trajectories in a dense road network due to the use of either the number of overlapping road segments or maximum/minimum distance between trajectories in their computation. In our work, we use the Dijkstra based dynamic time warping (DTW) distance measure, trajDTW \[33\] to compute trajectory similarities which is suitable for a large number of overlapping trajectories in a dense road network. The superiority of the trajDTW over the traditionally used dissimilarity with length (DSL) and Hausdorff distance measures is demonstrated in \[34\]. The trajDTW is a normal DTW algorithm with a Dijkstra distance matrix based cost function and a window parameter \( w \), which is set to the half of the length of shorter of two trajectories, to avoid overestimation of the actual distance. As the road network is static, the distance matrix \( D_{all} \) (of size \( |E| \times |E| \)) of all the edges \( E \in G_{RN} \) can be pre-computed and stored.

#### C. Non-directional trajDTW

The directionality of trajectories can result in misleading distances among them, which in turn may cause incorrect clustering results. For example, suppose there are two trajectories \( T_1 \) and \( T_2 \), which follow the same route but in opposite directions, then the distance between them considering their directions in computation will be higher than the distance computed without considering their directions. Therefore, if their movement direction is considered as part of the distance computation, \( T_1 \) and \( T_2 \) may not be grouped in the same cluster. The problem of incorrect distance measure due to the movement direction of trajectories is addressed by reversing one of them (reversing the sequence order so that the starting point becomes the ending point and vice versa), and taking the minimum distance between the first and second trajectory, and the first trajectory and second reverse trajectory. This distance is called non-directional trajDTW \[35\], and is given as:

\[
\text{non-directional trajDTW}(T_1, T_2) = \min(\text{trajDTW}(T_1, T_2), \text{trajDTW}(T_1, \text{Reverse}(T_2)))
\]

### Table I: Notations

| Symbol | Definition |
|--------|------------|
| \( \mathcal{F} \) | The set of trajectories |
| \( T_\ell \) | The \( \ell \)th trajectory of set \( \mathcal{F} \) |
| \( L \) | The length (or number of segments) of trajectory \( T \) |
| \( R_i \) | The \( i \)th segment of trajectory \( T_i \) |
| \( N_r, n \) | Number of trajectories in \( \mathcal{F} \) and MMRS sample \( S \), respectively |
| \( k, K \) | Number of non-directional and directional clusters in \( D \) |
| \( \mathcal{F}_j \) | Set of trajectories in cluster \( j \) |
| \( N_j \) | Number of trajectories in cluster \( j \) |
| \( \mathcal{R}_I \) | Set of points (segments) in cluster \( j \) |
| \( \mathcal{R}_S \) | Set of cluster of trajectories |
| \( R_{FRS}, R_{FSS} \) | Frequent road segment (FRS) and the set of FRSs, respectively |
| \( R_{SS}, R_{FSS} \) | Source segment and the set of SSs, respectively |
| \( M_j \) | Transition probability matrix for cluster \( j \) |
| \( W_j \) | Transition count matrix for cluster \( j \) |
D. The clusiVAT Algorithm

In our proposed framework, we modify the clusiVAT algorithm and use it for efficiently clustering large volumes of trajectory data. The clusiVAT model finds its root in the visual assessment of cluster tendency (VAT) [35] and improved VAT (iVAT) [36] algorithms. The VAT and iVAT algorithms reorder the input dissimilarity matrix \( D_N \) (for \( N \) datapoints) to \( D'_N \) (in VAT) and \( D'_N \) (in iVAT), respectively, using a modified Prim’s algorithm (for finding a minimum spanning tree (MST)) and by applying a graph theoretic distance conversion, such that the dark blocks along the diagonal of the reordered image \( I(D'_N) \) (or \( I(D'_N) \)) potentially represent different clusters. VAT and iVAT seem to work well for relatively small size datasets. However, both have space and time complexity of \( O(N^2) \), which limits their usefulness for input matrix sizes of an order of \( 10^3 \) and so. To overcome this limitation, an intelligent sampling based scalable clustering algorithm, clusiVAT [37] was proposed for visual cluster tendency assessment and subsequent clustering on big data. The essential steps in clusiVAT are:

Maximin Random Sampling (MMRS): MMRS sampling begins by finding \( k' \) (an overestimate of \( k \)) Maximin samples (distinguished objects), which are furthest from each other in the input data. Then, each object in the input data is grouped with its nearest Maximin sample with NPR. This step divides the entire data into \( k' \) groups. Then, a sample \( S \) of size \( n \) (just a small fraction of input data size, \( N \)) is built by selecting a proportional number of random data points (Random sampling (RS)) from each of the \( k' \) groups. Hence the term MMRS is used for the overall process. Any value of \( k' \) which overestimates assumed true number of clusters (k) i.e., \( k' \geq k \) should be a good choice [37]. For \( n, n = a few \) hundred datapoints is a reasonable choice for most datasets [38].

Step 2. iVAT: The iVAT is applied to the small \( D_n \) (computed from \( n \ll N \) samples) distance matrix to obtain its reordered distance matrix \( D'_n \). The image \( I(D'_n) \) usually provides an useful estimate of \( k \), without the need to compute the very large distance matrix \( D_N \).

Step 3. Clustering: Single linkage clusters are always aligned partitions in the VAT/iVAT reordered matrices. Having the estimate of \( k \) from the previous step, we cut the \( k - 1 \) longest edges in the iVAT-built MST of \( D_n \), resulting in \( k \) single linkage clusters. If the dataset is complex and clusters are intermixed, cutting the \( k - 1 \) longest edges may not always be a good strategy as the outliers, which are typically furthest from normal clusters, might comprise most of the \( k - 1 \) longest edges of the MST, resulting in misleading partitions. A useful approach in such a scenario is to manually select the dark blocks, and find the sample trajectories representing each dark block. Another useful approach [39] to obtain clusters is by cutting the MST using cut threshold magnitudes ordered by edge distances \( d \) in the MST. The cluster boundaries are defined by those indices \( z \), which satisfy

\[
d_z > \alpha \times \text{mean}(d),
\]

where \( \alpha \) is a parameter that controls how far two groups of data points should be from each other to be considered as separate clusters. Smaller values of \( \alpha \) represent tighter cluster boundaries, while large values of \( \alpha \) create loose cluster boundaries. The procedure to find an optimal value of \( \alpha \) is described in [39].

Step 4. Nearest Prototyping Rule (NPR): The \( k \)-partitions of the \( n \) samples are non-iteratively extended to the remaining (non-sampled) \( N - n \) objects in the dataset using the nearest prototyping rule.

The implementation and pseudocodes of the trajDTW, VAT, iVAT, and clusiVAT algorithms are well documented in the literature, and are not produced here for brevity.

E. Markov Chain Model

A Markov chain (MC) is the simplest form of the Markov process in which only the current state determines the probability of transitioning to the next state. Specifically, a Markov chain model is defined by the transition matrix \( M \), which contains the transition probabilities associated with various state changes. In a road network, an MC is constructed by assigning a state to each node or road segments in the given road network. For any two adjacent road segments \( R_i \) and \( R_j \) in road network \( G_{RN} \), the transition probability of traveling from \( R_i \) to \( R_j \) in one step is given by

\[
p_{ij} = \frac{\#(R_i, R_j)}{\#(R_i)},
\]

where \( \#(R_i, R_j) \) is the number of trajectories that contain the sequence \( \{R_i, R_j\} \) and \( \#(R_i) \) is the total number of trajectories that passes through \( R_i \). For each pair of adjacent road segments in the graph network, the transition probabilities can be computed using (4), and stored as entries \( M_{ij} \) of transition probability matrix \( M \) (of size \( |E| \times |E| \)). We also define a transition count matrix \( W \) whose \( i,j \)-th entry \( W_{ij} \) represents the number of trajectories that contain sequence \( \{R_i, R_j\} \) i.e., \( W_{ij} = \#(R_i, R_j) \). We utilize \( W \) in computing a representative trajectory for each cluster in our work.

IV. Proposed Framework

This section presents our proposed framework for trajectory prediction. The frequent route patterns of moving objects can be discovered by clustering their historical trajectories. In our framework, we employ a modified version of the clusiVAT algorithm that we called Traj-clusiVAT. In Traj-clusiVAT, we introduce a representative trajectory for each cluster to improve the performance of NPR for trajectory clustering. We also modify the NPR technique in Traj-clusiVAT to improve its performance for trajectory prediction. The Traj-clusiVAT algorithm partitions the trajectories into different groups of similar trajectories, based on the trajDTW distance measure. After clustering trajectories, we train a first-order Markov chain model for each cluster using only the trajectories contained therein. Then, these trained Markov chain models are used for trajectory prediction. The architecture of our proposed framework consisting of both training and prediction models is illustrated in Fig. [1].

A. Training Model

The essential steps of our training model are: (i) MMRS sampling on input trajectory data, (ii) VAT/iVAT and clustering
the trajectory sample using non-directional trajDTW to obtain k non-directional clusters (iii) VAT/iVAT and clustering the trajectories of each of the k clusters using trajDTW resulting in K (approx. 2k) directional clusters (iv) Compute representative trajectory (RT) of each cluster, (v) Assign remaining non-sampled trajectories to K clusters using NPR (vi) Re-compute the RT of each cluster, and (vii) Train a first-order Markov chain model for each cluster. The first six steps constitute the Traj-clusiVAT clustering algorithm. Below, we explain each step corresponding to the steps as shown in Fig. 1

1) MMRS sampling on input trajectory data

The first step consists of extracting a small, representative sample from the large trajectory data using MMRS sampling with non-directional trajDTW distance measure on input trajectory data \( T \). The aim of this step is to find the most distinguished vehicle routes in a given road network. The use of non-directional trajDTW circumvents the selection of more than one trajectory from the same route. In this way, the Maximin (first) step of MMRS ensures that MMRS samples contain the k' MM trajectories of the most distinguished vehicle routes. This divides the trajectory data \( T \) into k' partitions. Then, additional trajectories are randomly chosen from each of the k' partitions to generate a sample \( S \) of n trajectories. The MMRS intelligently chooses n trajectories which are almost equally distributed among the different clusters as the N trajectories in the big trajectory data, i.e., it obtains a representative sample.

2) Clustering trajectory samples using non-directional trajDTW

The previous step provides a trajectory sample S containing n trajectories. In this step, the iVAT is applied to the distance matrix \( D_n \) returning a reordered distance matrix \( D'_n \), and the cut magnitudes of the MST links, \( d \). The visualization of \( D'_n \) using \( I(D'_n) \) suggests the number of clusters \( k \) present in the dataset. The \( k \) partitions can be obtained by cutting the \( k - 1 \) edges or by cutting the MST cut magnitudes \( d \) using cut threshold \( \alpha \), as mentioned in Section [II.C]. The non-directional trajDTW distance measure is used in this step to cluster the n trajectories in order to avoid incorrect clustering due to the movement direction of trajectories, as mentioned in Section [II.C]. From here on, we denote \( k \) as the number of non-directional clusters.

3) Clustering trajectories in each cluster using trajDTW

The previous step clusters the trajectories based on their path similarity computed using non-directional trajDTW, which ensures that the trajectories that are in opposite directions, but follow similar routes, are clustered together. Since Markov chain models are used in our framework to model the trajectories of each cluster, their transition probabilities may be misleading for trajectory prediction task for clusters in which the number of trajectories in opposite directions is approximately equal. To circumvent this problem, we use the trajDTW (directional) distance measure for the sample trajectories of each cluster obtained in the previous step to separate the trajectories going in opposite directions using a second application of the iVAT algorithm, which in turn, gives \( K \sim 2k \) directional clusters.

4) Computing the RT of each cluster

In the NPR (next) step of clusiVAT, the non-sampled trajectories are assigned to one of the clusters (found in the previous step) based on their (nearest) distances from each cluster. For a fast and reliable implementation of NPR, we require a representative trajectory (RT) for each cluster that best describes the cluster, much like centroid-based clustering methods identify a representative "center" for each cluster. However, it is not possible to compute the centroid of trajectory clusters in a conventional way due to different lengths of trajectories in each cluster. Existing methods of calculating RT [40–43] in the literature either compute the mean trajectory using the average of GPS coordinates [41, 43]; or select a trajectory from each cluster which minimizes the dissimilarity between all the trajectories within the cluster [40, 42]; or pick a random trajectory [42] from each cluster, and designates it as the RT. These methods incur a large computational cost to compute an RT that minimizes the dissimilarity among all the trajectories. Additionally, RTs computed using these methods do not show all the possible variability inside a cluster [44]. The mean trajectory computed from trajectories of different lengths may be inaccurate; thus, it may not be a good representative of the cluster.

Our scheme generates an imaginary trajectory (IT) (it may not belong to any of the trajectories in the cluster) as an RT...
for each cluster that describes the major movement patterns of the trajectories belonging to that cluster. The pseudocode of our proposed method to compute RT for each cluster is shown in Algorithm 1. Below, we explain our RT computing algorithm.

First, we compute the transition count matrix $W_i$ for each cluster $T_j$ using the trajectories in that cluster (line 2). Then, for each cluster $T_j$, we compute the set of frequent road segments (FRSs) $\mathcal{R}_{FRS}^j$ using the $MinT$ threshold (line 3). The road segments in cluster $T_j$ which contains at least $MinT\%$ of the total trajectories in that cluster are assigned to $\mathcal{R}_{FRS}^j$. Then, a set of frequent source segments (FSSs) $\mathcal{R}_{FSS}^j$ is identified (line 4). A source segment $R_{FSS}^i$ is a FSS, $R_{FSS}^i$, if at least $MinT\%$ of total trajectories in the cluster originate from $R_{FSS}^i$, i.e., $R_{FSS}^i \in \mathcal{R}_{FSS}^j \subset \mathcal{R}_{FRS}^j$. Then for each FSS $R_{FSS}^i$, an imaginary trajectory $IT^i(R_{FSS}^j)$ is initialized with $R_{FSS}^i$, assigning it as current segment $R_{current}$ (lines 6–7). In lines 9–17, we compute the next RS $R_{next}$ based on the highest transition count from current RS $R_{current}$ using transition count matrix $W_i$ (refer to Section III-E). If $R_{next}$ is a FRS, then $R_{next}$ is added to current imaginary trajectory $IT^i(R_{FSS}^j)$, and assigned as $R_{current}$ to compute new $R_{next}$. The steps in lines 9–18 are repeated until $R_{next}$ is a non-FRS, which means an imaginary trajectory is an ordered sequence of only frequent road segments in that cluster. A total of $|\mathcal{R}_{FSS}^j|$ imaginary trajectories will be generated for each cluster $T_j$, corresponding to each $R_{FSS}^i \in \mathcal{R}_{FSS}^j$. We define a variable Count_score (line 8) for each imaginary trajectory $IT^i(R_{FSS}^j)$, $R_{FSS}^i \in \mathcal{R}_{FSS}^j$, which is the sum of the total transition counts of each RS $\in IT^i(R_{FSS}^j)$ in cluster $T_j$. Among all $|\mathcal{R}_{FSS}^j|$ ITs, the one which has the highest Count_score will be assigned as $RT(T_j)$ of cluster $T_j$ (line 20). As the RT($T_j$) is the sequence of FRS with highest Count_score, it contains major movement behaviour or patterns of the trajectories belonging to the cluster $T_j$.

Algorithm 1 does not require the computation of dissimilarity among all trajectories in that cluster to compute RT, which is computationally expensive for large size clusters. In contrast, Algorithm 1 is a novel algorithm to compute RT based on the transition count matrix of each cluster.

5) Assigning non-sampled trajectories to identified K clusters using NPR

The previous step gives representative trajectory $RT(T_j)$ for each cluster $T_j$. In this step, $N - n$ non-sampled trajectories are assigned to one of the $K$ directional clusters based on the NPR. The NPR method in clusiVAT uses the trajDTW (directional) distance measure to assign non-sampled trajectories to one of the $K$ clusters based on their nearest distance from (clustered)

Algorithm 1 Computing the RT of each cluster

Input: $\mathcal{T}_j$ - set of trajectories in cluster $j$, $N_j$ - number of trajectories in cluster $j$, $\mathcal{R}_j$ - set of road segments in cluster $j$, $\mathcal{C}(\mathcal{T}) = \{T^1, ..., T^K\}$ - set of cluster of trajectories, MinT- FRS threshold

1: for each cluster $T_j \in \mathcal{C}(\mathcal{T})$ do
2: Compute FRSs, $\mathcal{R}_{FRS}^j$, from $\mathcal{R}_j$, $\mathcal{R}_{FRS}^j = \{R_j \in \mathcal{R}_j | MinT \times N_j \}$
3: Compute FSSs, $\mathcal{R}_{FSS}^j$ from $\mathcal{R}_{FSS}^j = \{R_i \in \mathcal{R}_{FRS}^j \}$
4: for each FSS $R_{FSS}^i \in \mathcal{R}_{FSS}^j$ do
5: Assign $R_{FSS}^i$ as current road segment, $R_{current} = R_{FSS}^i$
6: Initialize an imaginary trajectory $IT^j$ with $R_{current}$, $IT^j(R_{FSS}^i) = \{R_{current}\}$
7: Count_score($IT^j(R_{FSS}^i)$) = 0
8: while each RS of $IT^j(R_{FSS}^i) \in \mathcal{R}_{FRS}^j$ do
9: Compute next RS, $R_{next} = \arg\max_{R_i \in \mathcal{R}_j} \{W_i\}$
10: if $R_{next} \in \mathcal{R}_{FRS}^j$ then
11: Append $R_{next}$ to existing $IT^j(R_{FSS}^i)$
12: $R_{current} = R_{next}$
13: Count_score($IT^j(R_{FSS}^i)$) += $W_i$
14: else
15: break;
16: end if
17: end while
18: end for
19: Select $IT^j(R_{FSS}^i)$ with the highest Count_score($IT^j(R_{FSS}^i)$) from all $|\mathcal{R}_{FSS}^j|$ ITs of $T_j$, and assign it as RT for cluster $T_j$
20: end for

Output: $RT(T_j)$ - RT for each cluster $T_j \in \mathcal{C}(\mathcal{T})$

Algorithm 2 Hybrid NPR Method

Input: $T_q$ - query trajectory, $M_i$ - transition probability matrix for cluster $j$, $RT(T_j)$ - representative trajectory for cluster $j$

1: Compute the path probability $P(T_q)$ of query trajectory in each cluster $T_j$ using $M_i$ and Eq. 5
2: if any($P(T_q) > 0$) then
3: Select the cluster $c$ with the highest $P(T_q)$ i.e., $c = \arg\max_{T_j \in \mathcal{C}(\mathcal{T})} \{P(T_q)\}$
4: else
5: Compute the trajDTW distance of $T_q$ from $RT(T_j)$, $y = \text{trajDTW}(T_q, RT(T_j))$, for each cluster $T_j \in \mathcal{C}(\mathcal{T})$
6: Select the cluster $c$ with the minimum $y_j$ i.e., $c = \arg\min_{T_j \in \mathcal{C}(\mathcal{T})} \{y_j\}$
7: end if
8: Assign the $T_q$ with cluster $c$ (or $T_j$).

Output: cluster label for $T_q$
sample trajectories. However, trajDTW distance of a non-sampled trajectory to cluster RTs may not be an appropriate measure for the NPR step due to its dependency on the length of trajectories, as explained by the following example.

Suppose $T_a$ is a non-sampled trajectory in $T$, and $RT(3')$ and $RT(3')$ are the RT of cluster $3'$ and $3'$, respectively. Let $T_b$ be a sub-trajectory of $RT(3')$ i.e., $T''$ is fully contained in $RT(3')$. Since the trajDTW distance relies on a warping window size parameter $w$, the $trajDTW(T_a, RT(3'))$ not only depends on the coordinates of RSs of both trajectories, but it also depends on the length of $T_a$ and $RT(3')$. Moreover, $trajDTW(T_b, RT(3'))$ also varies depending on the position of $T_a$ in $RT(3')$ due to window parameter. Therefore, even if $T_a \subseteq RT(3')$ and $T_a / RT(3')$, $T_a$ may be incorrectly assigned to cluster $3'$ instead of $3'$ if $\text{trajDTW}(T_a, RT(3')) \geq \text{trajDTW}(T_b, RT(3'))$. Here is such an example from T-Drive data. Suppose $T_1 = \langle 70, 75, 90, 89, 88 \rangle$ is a non-sample trajectory, and $RT(3') = \langle 16, 18, 68, 70, 75, 90, 89, 88 \rangle$ and $RT(3') = \langle 68, 70, 75, 91, 92 \rangle$ are RTs of two clusters, where each trajectory is represented by a sequence of road segments’ IDs of Beijing road network (Refer to Section VI-A). The trajDTW distances are: $\text{trajDTW}(T_1, RT(3')) = 0.3482$ and $\text{trajDTW}(T_1, RT(3')) = 0.2767$. Therefore, although $T_1$ is a sub-trajectory of $RT(3')$, it will be assigned to cluster $3'$ based on nearest trajDTW distance. Such assignments of non-sampled trajectories to (incorrect) cluster may include outlier trajectories or road segments in that cluster, which may adversely affect Markov chain modeling, and consequently, degrade the performance of trajectory prediction.

To address above issue, we propose a hybrid NPR strategy based on the path similarity and trajDTW distance measure. Hybrid NPR is similar to clusiVAT NPR except for those non-sampled trajectories, which are sub-trajectory of any of the clusters’ trajectories. The pseudocode of our hybrid NPR method is shown in Algorithm 2. For a query trajectory $T_q = \{R_1, R_2, ..., R_l\}$, we first compute the path probability $P(T^q)$ for each cluster $3'$, which is defined as

$$P(T^q) = \prod_{j=1}^{l} P_j(j+1) \Leftrightarrow \prod_{j=1}^{l} M_{j(j+1)}^i.$$  

$P(T^q) > 0$ means that sequence $T^q$ appears at least once in cluster $3'$, whereas $P(T^q) = 0$ means that sequence $T^q$ is not present in cluster $3'$. If the sequence $T^q$ is present in any cluster $3'$ i.e., $\forall j \in \{N\}, (P(T^q)) > 0$, then $T^q$ is assigned to the cluster with the highest path probability. If the sequence $T^q$ is not present in all clusters $3'$ i.e., $\forall j \in \{N\}, (P(T^q)) = 0$, then $T^q$ is assigned to the cluster based on its (minimum) trajDTW distance from RTs. All non-sampled trajectories in $T$ are assigned to one of the $K$ clusters using Algorithm 2.

6) Recompute the RT of each cluster after NPR

The assignment of all non-sampled trajectories to one of the $K$ clusters in the NPR step updates each cluster with new trajectories. Therefore, a representative trajectory is recomputed for each updated cluster using Algorithm 3.

7) Train Markov chain model

For each of the $K$ clusters, we build a first-order Markov chain model using the trajectories of that cluster. Specifically, we compute the transition probability matrix $M'$ for each cluster $3'$.

For a basic understanding of Traj-clusiVAT algorithm, we graphically explain its steps on a small trajectory data $T$, as shown in Fig 2. An input trajectory data $T$ containing $N = 9$ trajectories is shown in Fig 2(a). The MMRS sampling on $T$ with non-directional trajDTW in the first step returns a MMRS sample $S$ containing $n = 6$ sample trajectories $\{1, 4, 5, 6, 7, 9\}$, which are well-distributed in sample $S$, as shown in Fig 2(b). In the next step, iVAT is applied to $S$ using the non-directional trajDTW distance measure, which clusters the trajectories based on the path similarity irrespective of their movement directions. The iVAT image in Fig 2(c) shows four dark blocks along its diagonal, which indicates four clusters in sample $S$. Having an estimate of $k = 4$, sample $S$ is partitioned into four (non-directional) clusters $\{\{1, 4\}, \{5\}, \{6, 7\}, \{9\}\}$, as shown with four different colors in Fig 2(c). Then, the trajectories in each cluster going in opposite directions are separated using the iVAT with the trajDTW distance measure, which gives $K = 6$ directional clusters $\{\{1\}, \{4\}, \{5\}, \{6, 7\}, \{9\}\}$, each cluster is shown with a different colour in Fig 2(d). Since there are only one trajectory in each cluster in this case, they are the RTs for corresponding clusters. In the next step, non-sampled trajectories $\{2, 3, 8\}$ are assigned to one of the 6 clusters using NPR (Algorithm 2), which partitions the complete data into 6 clusters $\{\{1, 2, 3\}, \{4\}, \{5\}, \{6, 8\}, \{7\}, \{9\}\}$. Trajectory 4 is in different cluster than $\{1, 2, 3\}$ due to opposite direction. Then, a Markov chain model is trained for each cluster using the trajectories of that cluster.

B. Prediction Model

For a given partial trajectory $T^p = \{L_1, L_2, ..., L_m\}$, we first estimate the best matching representative cluster $3'^c$ using our hybrid NPR approach, and then choose the corresponding Markov model of the cluster to predict the next locations $L_i$, $i \geq m + 1$. Using the cluster $3'^c$, the location $L_{m+1}$ that the object will arrive at next is given by

$$L_{m+1} = \arg \max_{L_j \in \mathbb{R}^c} \{p_m\} \Leftrightarrow \arg \max_{L_j \in \mathbb{R}^c} \{M'_{mj}\}.$$  

The $T^p$ is updated with the next predicted location $L_{m+1}$. Then, the updated $T^p$ is used to estimate the best matching cluster and the corresponding MM is used to predict the next location. The complete trajectory is predicted by computing next locations in a sequential manner using these steps.

V. TIME COMPLEXITY

In this section, we discuss the time complexity of our proposed Traj-clusiVAT based TP approach. The first step in Traj-clusiVAT is the selection of $k'$ distinguished trajectories which are at maximum distance from each other. This step has the time complexity of $O(k'N)$, where $k'$ is a user-defined parameter for an overestimate of the number of clusters in the input trajectory data and is usually chosen to be (inessentially) large (usually 50 to 200). The next step is to randomly select $n$ trajectories from $k'$ NPR groups to get a sample $S$. The computation of distance matrix $D_{n}$ and VAT on a sample $S$ has a time complexity of $O(n^2)$. Usually $n << N$, so the
computations of $D_a$ and VAT on $S$ is pretty fast and takes just a small fraction of the total run time of Traj-clusiVAT. The trajDTW distance measure uses Dijkstra’s shortest path distance in the standard DTW algorithm. Its best, average case complexity with binary heaps is $O(|E|+|V| \log |V|)$ [45]. For two trajectories of length $l_1$ and $l_2$, standard DTW has time complexity of $O(l_1l_2)$. Remark- There are approximate algorithms such as FastDTW [46] which have a linear time complexity in the average length of trajectories, however, we have not used this implementation in our experiments. The NPR step in Traj-clusiVAT has complexity of $O(n(N-n))$. The computation of RT’s have linear time complexity in $K$. The construction of a Markov model for each cluster is a simple and fast process, which has $O(K)$ time complexity and $O(|E|^2)$ space complexity.

VI. EXPERIMENTS

In this section, we conduct an extensive experimental study on two real-life, vehicle trajectory datasets to evaluate the performance of our proposed framework. We first describe the datasets, their preprocessing, evaluation metrics and computational protocols adopted in our empirical study, and then present the experimental results.

A. Datasets

We performed our experiments on two real trajectory datasets.

1) T-Drive taxi trajectory data [47]

This trajectory dataset is obtained from the T-Drive project which contains one-week trajectories of 10,357 taxis during the period of Feb. 2 to Feb 8, 2008 within Beijing, China. The total number of points is about 15 million and the total distance of the trajectories is 9 million kilometres. In our experiment, we have taken a subset of this dataset, which contains trajectories from a road network in the center of Beijing city, as shown in Fig. 3(a). This road network consists of 100 nodes and 141 road segments (edges). The average sampling interval is 177 seconds with an average distance of about 623 meters, which is quite large for a city traffic environment as the length of many road segments is smaller than the average sampling distance.

2) Singapore taxi trajectory data

This dataset consists of the trajectories of more than 15,000 taxis collected over a duration of 1 month from a road network in Singapore City, as shown in Fig. 3(b). This dataset is very dense as it consists of more than 370 million GPS logs. The general format of each data point is as follows: [Time Stamp, Taxi Registration, Latitude, Longitude, Speed, Status]. The Status field contains information about the occupation state of Taxi, such as FREE and POB (Passenger on Board). In order to extract each individual taxi’s trip from the raw data, we detect the following sequence: starting from FREE to POB and ending from POB to FREE, using the trip extraction framework presented in [48]. This road network consists of 1641 nodes and 2941 edges, with an average edge length of 350m.

Data Pre-processing

To obtain the trajectories as a sequence of road segments, we use the popular open source map matching tool GraphHopper [49], which provides an implementation of the approach presented in [50].

After pre-processing, we have $N = 43,405$ trajectories in the T-Drive data whose lengths lie in the range of 5 to 200 road segments and have an average of 14 road segments, and $N = 3,259,290$ (3.26 million) trajectories in the Singapore data whose lengths lie in the range of 10 to 250 road segments and have an average of 22 road segments. To prepare training and test sets for both datasets, we first divided the trajectories into two sets based on the day of week viz., weekdays and weekends, during which the trip is being made. For the one-week T-Drive data, we considered trajectories during first 4 weekdays (Monday to Thursday) and first weekend day (Saturday) as the training set, and trajectories during the remaining days (Friday and Sunday) of that week as the test set. For the one-month Singapore data, we considered 60% trajectories randomly as training set and remaining 40% as the test set, for both weekdays and weekend data. The size of training and test sets for both trajectory datasets is shown in Table II. We split each trajectory in a test set into two halves. The first half is used as a partial trajectory (or query trajectory) for predicting its future locations and the second half is used as ground truth to validate our predictions. The distribution of predicted trajectories (second half) in the T-Drive and Singapore test sets is shown in Fig. 4.

B. Evaluation Metrics

In our experiments, we assess the performance of our framework for next location prediction (also known as one-
step prediction) and long-route prediction using following evaluation metrics:

1) **Prediction Accuracy (PA)**

The PA is the ratio of correctly predicted locations to the total possible number of predicted locations for each trajectory. Given a predicted trajectory sequence $T_{pred} = \{L_1, L_2, ..., L_m\}$ and a true (actual) trajectory sequence $T_{true} = \{R_1, R_2, ..., R_m\}$, the prediction accuracy is defined as

$$PA = \frac{1}{|T_{pred}|} \sum_{j=1}^{m} H(L_j, R_j),$$

where $H(L_j, R_j) = 1$ if $L_j = R_j$, else 0. The average prediction accuracy is the average of PA for all predicted trajectories in test set $T_{test}$.

2) **Prediction Rate (PR)**

The PR is the number of trajectories that are correctly predicted over the total number of trajectories in test set. It is defined as

$$PR = \frac{1}{|T_{true}|} \sum_{j=1}^{m} H(T_{pred}, T_{true}),$$

where $H(T_{pred}, T_{true}) = 1$ if $T_{pred} = T_{true}$, else it is 0.

3) **Distance error (DE)**

Another important performance metric of the long-term prediction system is the capability of continuous route prediction. The distance error is defined as the average spatial (Haversine) distance between predicted and actual routes. Given a route sequence $T_{pred}$ and $T_{true}$, the distance error between them is given as

$$DE(T_{pred}, T_{true}) = \frac{1}{|T_{pred}|} \sum_{j=1}^{m} D_H(L_j, R_j),$$

where $D_H(L_j, R_j)$ is the Haversine [51] distance between two locations (road segments).

4) **One-step accuracy (OA)**

This is the ratio of correctly predicted next locations to the total predicted next locations for all trajectories in test set.

5) **One-step distance error (ODE)**

The ODE defined as the average distance error for one-step (or next location) prediction.

C. **Comparison Methods**

Among the plethora of MM and clustering based TP methods available in the literature, we implemented these two approaches for comparison.

1) **Mixed Markov model (MMM) based TP** [8]: MMM was proposed as an intermediate model between standard MM and HMM which can encompass all types of movement behaviour present in an input trajectory data. It first clusters the trajectories into groups using the EM algorithm, and then builds an MM for each group, which is subsequently used for prediction. This approach was tested on synthetic and real datasets in [8], which showed 74.1% accuracy for MMM, in comparison to 16.9 – 45.6% for MM and 2.4 – 4.2% for HMM.

2) **NETSCAN-based TP**: The well-known density-based algorithm DBSCAN and its variants [32], [52] – [54] have been used extensively as a trajectory clustering method for location prediction [11]. However, they are not suitable for a large number of trajectories as computation of the distance matrix is time intensive. Kharrat et al. [9] proposed a trajectory clustering relative of DBSCAN, called NETSCAN which first finds dense road segments based on the moving object counts, merges them to form dense paths on the road network, and then assigns sub-trajectories to the dense paths based on a measure of similarity. This method requires two user-defined parameters: a density threshold – the minimal required density for transition, and a similarity threshold – the maximum density difference between neighbouring road segments. We implement NETSCAN to cluster trajectories into dense road segments, then built an MM for each cluster, and subsequently used them for TP.

Our proposed method and the baseline methods discussed above are also comparable in terms of prediction time (which will be discussed shortly). They all require a short prediction time and satisfy the requirement of real-time prediction.

D. **Computation Protocols**

All algorithms were coded in MATLAB on a PC with the following configuration; OS: Windows 7 (64 bit); processor: Intel Core i7 – 4770 @3.40GHz; RAM: 16GB. We denote the comparison approaches of [8] as MMM, of [9] as NETSCAN, and our Traj-clusiVAT based TP approach as TrajclusiVAT. All three algorithms were applied to T-Drive data. The MMM method requires the computation and storage of an intermediate matrix of size $|E| \times |E| \times N$, which is very large for Singapore data (due to large $|E|$ and $N$), so we can not apply MMM to the Singapore data. However, we have compared it with NETSCAN-TP and Traj-clusiVAT-TP on a subset obtained from a smaller part of the Singapore road network. The number of mixed models of MMM was determined using 10-fold cross-validation. The NETSCAN parameter, density threshold and similarity threshold, were chosen to get as many dense paths (with at least six road segments) as the number of clusters we get using the TrajclusiVAT algorithm, for a fair comparison. The parameters for Traj-clusiVAT were chosen as follows: $k' = 150$, $n = 500$, and $\alpha = 0.05$ for the T-drive data, and $k' = 300$, $n = 1000$, and $\alpha = 0.06$ for the Singapore data, and $Min T = 30\%$ for both data. It is worth noting that, unlike other clustering algorithms, the clusiVAT algorithm is relatively insensitive to the choice of $k$ and $n$ [5]. Moreover, we study the effect of $\alpha$ on Traj-clusiVAT performance in our experiments.

E. **Comparison of MMM, NETSCAN, and Traj-clusiVAT for Long-term predictions**

Long-term prediction, also known as continuous route prediction, is a challenging and ongoing research problem in TP. In this experiment, we compare the performance of the MMM, NETSCAN, and Traj-clusiVAT-based prediction approaches for $m$-step predictions. Specifically, this refers to predicting the next $m$ locations for a given partial trajectory. Fig.[5] shows the average prediction accuracy (left panels) and average distance
error (right panels) of all three algorithms for increasing prediction steps. The graphs in Fig. 5 support these observations:

(i) First, the Traj-clusiVAT outperforms the MMM and NETSCAN-based TP approaches based on the average PA and DE for the T-Drive data, as shown in Fig. 5(a). The higher the number of prediction steps, the larger the gap between Traj-clusiVAT and other two approaches. This means that the Traj-clusiVAT performs better not only for short-term predictions but it performs even better than the other two approaches for long-term predictions. This is probably because Maximin sampling in Traj-clusiVAT finds the trajectories which are furthest from each other. As the trajDTW distance measure yields higher distances for longer trajectories, Maximin sampling tends to pick longer trajectories in its output sample which form separate clusters in subsequent steps. The Markov models trained on these clusters after the NPR step contain all movement behaviours similar to those longer trajectory patterns. Therefore, if a query trajectory pattern is not available in any cluster, which is frequent for longer query patterns, then it is assigned to a cluster based on its nearest distance from all cluster RTs. This will assign longer query trajectories to any of the clusters containing longer trajectory patterns, and subsequently, corresponding MMs trained on these clusters contribute towards better predictions for longer query trajectories during the prediction phase. On the other hand, the longer movement rules cannot be easily represented by Markov-based models, especially for irregular trajectory data, due to uncertainty in movement behaviours of vehicles in a complex road network. As there are only a few prediction trajectories available for the T-Drive test set whose lengths are greater than 16 as shown in Fig. 4(a), the performance of all approaches cannot be considered conclusive for longer prediction steps ($m > 16$) based on their performance on the T-drive data.

(ii) Fig. 5(b) shows that the Traj-clusiVAT model also performs better than the NETSCAN-based method based on the average PA and DE values for the Singapore data. The gap between the NETSCAN and Traj-clusiVAT plots increases until 31th prediction step and then reduces with longer prediction steps. This may be because the trajectory clusters obtained by NETSCAN are usually spread over the entire road network, which results in longer dense paths. Therefore, its performance becomes competitive with Traj-clusiVAT for longer prediction lengths compared to its short-term prediction performance.

(iii) It can be observed that difference in performance of the NETSCAN method and the proposed method is less for short-term and more for long-term prediction for T-drive data, whereas it is opposite for the Singapore data. This is because the T-drive dataset contains many parallel and perpendicular road segments and intersections that span only a small road network (Beijing city center), whereas, Singapore dataset contains relatively longer and straight (fewer intersections) trajectories (compared to T-Drive subset) that span entire Singapore city road network (significantly bigger than Beijing city center road network). Therefore, incorrect predictions cause relatively smaller distance error for T-Drive data as compared to distance error for Singapore taxi dataset. NETSCAN’s clusters for Singapore dataset are ordered sequences of only a few but longer and straight dense paths, therefore, it modeled long trajectories better for Singapore dataset, and in turn, performed better for long-term prediction as compared to the T-Drive dataset.

(iv) The performance of all three approaches deteriorates as the prediction step increases. This may be because the number of frequent trajectory patterns obtained is small for long-term predictions, which do not contain enough information to forecast future locations.

1And the other reason, as Niels Bohr said, is that “it is very hard to predict, especially the future”
TABLE III: Long-term prediction: Comparison of MMM, NETSCAN and Traj-clusiVAT

| T-Drive Data | MMM | Average PA | Average DE (km) | PR (%) |
|--------------|-----|------------|----------------|--------|
| NETSCAN      | 0.41| 0.87       | 39.9           |
| Traj-clusiVAT| 0.62| 0.58       | 49.8           |
| Singapore taxi data | NETSCAN | 0.34 | 1.41 | 5.1 |
|               | Traj-clusiVAT | 0.59 | 0.60 | 24.8 |
| Singapore taxi subset | MMM | 0.59 | 0.30 | 32 |
|               | NETSCAN      | 0.54 | 0.33 | 19 |
|               | Traj-clusiVAT| 0.64 | 0.25 | 46 |

In our experiments, we find that most of the clusters contain frequent trajectory patterns whose lengths are less than six or seven. Only a few clusters contain frequent trajectory patterns whose lengths are longer than seven steps. This finding conforms with the real-world situation, where a driver usually predicts only next few locations.

The average long-term prediction performance of all three approaches is summarized in Table III. The best performance is shown in bold for both datasets. Traj-clusiVAT achieves the highest PA, 0.62 and 0.59 and the lowest DE, 0.58 km and 0.60 km, for the T-Drive and Singapore taxi datasets, respectively. The MMM-based prediction approach is the second best method for T-Drive in terms of all three evaluation metrics. Traj-clusiVAT achieves prediction rates of 49.8% and 24.8% for the T-Drive and Singapore trajectory datasets, respectively. In other words, Traj-clusiVAT is able to predict complete trips for around 50% of the trajectories in T-Drive, and for around 25% of the trajectories in Singapore data. In contrast, MMM predicts about 40% of the total trajectories correctly for the T-Drive dataset. Although, NETSCAN performance improved for longer predictions due to longer dense paths, it only predicted about 5% of the total trajectories correctly. Overall, Traj-clusiVAT based prediction approach outperforms both MMM and NETSCAN-based prediction approaches based on all three evaluation metrics.

As we could not apply MMM to the Singapore data due to its high computation and space complexity, we applied it to a subset of a small part of the Singapore road network, as shown in Fig. 6 (a). This sub-graph consists of 238 nodes, 417 edges, and 828,870 trajectories. Similar to our previous experiments, we considered 60% of the trajectories (497,320) randomly as training data and the remaining 40% (331,550) as the test set, for both weekdays and weekend data. Although we could reduce the computational time and space requirements by considering a smaller road network, it still has high complexity due to large $N$ (497,320). We could not run MMM with more than 10,000 trajectories as it started to slow down our PC significantly and gave "Out of Memory" error message. Therefore, we used 10,000 randomly selected trajectories as a training set for MMM-based TP to avoid associated computational and storage overload. We used the same subset for NETSCAN-TP and Traj-clusiVAT-TP for training in this experiment.

Figs. 6 (b) (c) show the average prediction accuracy and average distance error, respectively, for all three algorithms. Traj-clusiVAT-TP method outperforms the MMM-based and NETSCAN-based TP approaches for this subset of the Singapore dataset, based on the average PA and DE. Overall, Traj-clusiVAT-TP achieves the highest average PA (0.64), lowest average DE (0.25 km) and highest PR (46%) among all three methods for Singapore taxi subset, as shown in Table III.

F. Next location predictions

In this experiment, we compare Traj-clusiVAT to the other two comparison approaches for predicting next locations. Given a taxi’s current location, the next location prediction is to forecast the next location where the taxi may go. Table IV shows the one-step accuracy (OA) and one-step distance error (ODE) on the T-Drive and Singapore trajectory datasets. The Traj-clusiVAT-based approach predicts next location with more than 80% accuracy and with distance error of less than a quarter of km for both T-Drive and Singapore data. The long-term prediction performance (Table III) of NETSCAN and Traj-clusiVAT is better for T-Drive than for the Singapore data. Conversely, the next location prediction performance of both approaches is better for the Singapore data than the T-Drive data. This may be because Singapore data contains a large number of longer trajectories that span entire Singapore city, whereas T-drive contains partial trajectories belonging to small part of the entire road network, hence modeling is not that efficient for T-drive data. In summary, Traj-clusiVAT outperforms both MMM and NETSCAN for next location prediction.
G. Effect of latest locations of partial trajectory for prediction

In the prediction step of Traj-clusiVAT, a partial trajectory $T^p = \{R_1, R_2, ..., R_l\}$ is assigned to one of the $K$ clusters using our hybrid NPR approach. For a $T^p$, the best cluster is chosen based on either its path probability $P^i(T^p)$ in each cluster or its trajDTW distance from each cluster (if $T^p$ is not fully contained in any cluster). The length of known partial trajectory $T^p$ increases after each next location prediction as $T^p$ is updated with a predicted location after each prediction, and subsequently, the updated $T^p$ is used for next location prediction, and so on.

We conduct an experiment in which instead of using full known partial trajectory $T^p$, we use only the latest movement steps or latest subsequence of $T^p$ until prediction to choose the best matching cluster in the hybrid NPR step. In this regard, we choose a different number of latest locations of known partial trajectories until prediction and investigate the effect on the performance for trajectory prediction.

Fig 7 shows the average distance error for a different number of latest locations of known partial trajectories until prediction for the T-drive and Singapore data. It can be inferred from the figure that the best performance is achieved when only the latest two or three locations of partial trajectory are used to find the best matching cluster. The average distance error increases if more than three latest locations are used to find the best cluster in the hybrid NPR step. This is because as the length of $T^p$ increases, its path probability in each cluster decreases, which means that the chance of sequence $T^p$ being fully contained in any cluster decreases. Moreover, if $T^p$ is not fully contained in any cluster representative trajectory, its distance from all the clusters increases with increasing length. This may result in wrong cluster assignment, which in turn, may degrade Traj-clusiVAT’s prediction performance.

H. Effect of Cut threshold $\alpha$

In this experiment, we study the effect of cut threshold $\alpha$. The parameter $\alpha$ in Traj-clusiVAT controls how far two groups of data points should be from each other to be considered as different clusters. Figure 8 shows the average DE and the number of clusters $K$ for different values of $\alpha$ for the T-Drive and Singapore data. The lower the cut threshold, the tighter the cluster boundaries, and hence, the higher the number of clusters. As the number of clusters $K$ increases, the average DE reduces. This is primarily because the higher $K$ corresponds to a larger number of unique frequent patterns, which improves the prediction performance. Figure 8 shows that the Traj-clusiVAT performance improves with lower cut threshold $\alpha$ or with the higher number of clusters. However, with a large $K$, more MM needs to be trained, and hence, system complexity increases. Moreover, Traj-clusiVAT performance does not improve significantly below a certain value of $\alpha$ for either dataset. The procedure to find an optimal value of $\alpha$ is described in [39].

I. Time performance analysis

The training time of all three algorithms on different-size training sets is shown in Fig. 9. The CPU-time for MMM increases most with the training data size because the computation of an intermediate matrix of size $|E| \times |E| \times N$ incurs high computational overhead and space complexity for large $N$. On the other hand, NETSCAN incurs the lowest computation time among all three methods. This is because it just computes dense paths based on the movement counts and density threshold, and assigns all trajectories to these dense paths based on similarity. Although it takes less time for training, it suffers from lower prediction accuracy. Traj-clusiVAT scales almost linearly in the number of trajectories, which make it scalable for big trajectory datasets.

![Fig. 7: Average DE vs latest locations of partial trajectory used to select best cluster in the hybrid NPR step.](image)

![Fig. 8: Effect of cut threshold $\alpha$](image)

![Fig. 9: Training time comparison](image)

Prediction-time is also an important criterion in real-time trajectory prediction. The average prediction time for all three approaches is presented in Table V. We can see that all three approaches take similar times to forecast each trajectory for the T-drive dataset. The response time is less than 1.5ms for T-drive, which suggests that all three approaches satisfy the requirement for real-time prediction. The average prediction time is higher for the Singapore dataset due to a large number of clusters identified by both NETSCAN and Traj-clusiVAT algorithms, but at $\sim 0.06$ seconds, it is negligible in terms of real-time prediction utility.

| TABLE V: Prediction time in seconds for all three algorithms |
|-------------------------------------------------------------|
| T-Drive Taxi | MMM | NETSCAN | Traj-clusiVAT |
|---------------|-----|---------|---------------|
| Singapore Taxi | 0.0014s | 0.0011s | 0.0012s |
| Singapore Taxi | 0.063s | 0.066s |
VII. CONCLUSIONS

Most existing TP approaches are not suitable for large volumes of overlapping trajectories in a dense road network. This article presents a novel, scalable, hybrid architecture for short-term and long-term trajectory prediction, which can handle a large number of trajectories from a large-scale dense road network. The proposed framework is based on a scalable clustering approach, called Traj-clusiVAT, which is a modified version of clusiVAT for trajectory prediction. In particular, Traj-clusiVAT develops a novel algorithm to compute a representative trajectory for each cluster. We also presented a new, hybrid nearest prototyping approach for accurate trajectory assignment to (one of) the clusters identified in previous steps of Traj-clusiVAT. Finally, we also propose a hybrid prediction framework based on hybrid NPR which can assign a query trajectory to best-matching cluster in a robust way to improve prediction performance.

We demonstrated the superiority of our proposed approach by comparing it with mixed Markov model and NETSCAN based TP approaches on two real trajectory datasets, including a large-scale trajectory dataset containing 3.28 million trajectories of passenger trips obtained from 15,061 taxis within Singapore over a period of one month. Our experimental results on both trajectory datasets show that Traj-clusiVAT based TP approach outperforms the other two approaches based on the prediction accuracy and distance error for short-term and long-term prediction for these two datasets. Our experimental results also suggest that Traj-clusiVAT satisfies the requirement for real-time predictions. Our next effort will focus on online training in Traj-clusiVAT using incremen- tal/decremental VAT approaches to update clusters in real-time. We also intend to include additional factors such as speed, time, and user information in our prediction system to improve its prediction performance.

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