# ON BOOSTING SPATIAL COMPUTATIONS FOR LOCATION-BASED SERVICES

### by

# CHENG LONG

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 $\mathbf{b}\mathbf{y}$ 

### CHENG LONG

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DR. RAYMOND CHI-WING WONG, THESIS SUPERVISOR

\_\_\_\_\_

PROF. QIANG YANG, HEAD OF DEPARTMENT

Department of Computer Science and Engineering
20 April 2015

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### CHENG LONG

Department of Computer Science and Engineering

The Hong Kong University of Science and Technology

### **ABSTRACT**

Nowadays, *location-based services* (LBSs), which refer to those services that are based on *location* (or *spatial*) data, are broadly used in our daily life. Some popular types of LBS include "search-nearby" which searches objects (e.g., restaurants, hotels and shops) near a location, "spatial crowdsourcing" which allows people to post tasks to be performed at a location (these people are called "requesters") and people to pick some tasks to perform (these people are called "workers"), and "trace tracking" which records the trace of a movement (e.g., the moving trace of a hiker). Each type of LBS usually relies on some computation based on spatial data (which is termed as *spatial computation*). For example, the "search-nearby" service relies on *spatial keyword query* to find all objects that are near a given query location and contain a given query keyword, the "spatial crowdsourcing" service relies on *spatial matching* to match between tasks and workers, and the "trace tracking" service relies on *trajectory data* 

#### management.

In this thesis, we introduce three techniques for boosting the spatial computations that are central to LBSs, namely the *collective spatial keyword query* which is one type of spatial keyword query and finds a set of spatial objects that cover all the given query keywords and have the smallest distance from the query location, *worst-case optimized spatial matching* which matches between two sets of spatial objects with the smallest worst-case cost, and *direction-preserving trajectory* which simplifies the trajectory while preserving the direction information embedded in the trajectory data.

### CHAPTER 1

### INTRODUCTION

Nowadays, *location-based services* (LBSs), which refer to those services that are based on *location* (or *spatial*) data, are broadly used in our daily life. Some types of LBSs include the "search-nearby" service which searches some places (e.g., restaurants, hotels and shops) near a location, the "spatial crowdsourcing" service which allows people to post tasks to be performed at a location, and the "trace tracking" service which records the trace of a movement.

Many apps for smart phones provide some types of LBS. Some examples that provide the "search-nearby" service include Yelp, Gowalla, Google Places, NearbyFeed, Sonar, OpenTable, and WeChat. Some examples that provide the "spatial crowdsourcing" service include LocalHands, Airtasker, gMission and iRain. Some examples that provide the "trace tracking" service include GeoFlyer, Maps+, Nike+ and Run Keeper. According to *appcrawlr.com*, there are more than 500 apps that provide LBSs.

Each type of LBS relies on some computation based on spatial data which is called *spatial computation*. For example, one type of spatial computation that is the core of the "search-nearby" service is *spatial keyword query*, which finds the nearest objects that contain a given query keyword to a query location, one for the "spatial crowd-sourcing" service is *spatial matching*, which finds a matching between two sets of spatial objects (since spatial crowdsourcing usually involves a procedure of matching the tasks with the workers), and one for the "trace tacking" service is *trajectory data management* (since the trace of a movement is usually represented by a trajectory).

In this thesis, we introduce three techniques, namely the Collective Spatial Key-

word Query which is one type of spatial keyword queries, Worst-Case Optimized Spatial Matching which is one type of spatial matching and Direction-Preserving Trajectory Simplification which could be used for trajectory data management.

Collective Spatial Keyword Query. Recently, spatial keyword queries become a hot topic in the literature. One example of these queries is the collective spatial keyword query (CoSKQ) which is to find a set of objects in the database such that it covers a set of given keywords collectively and has the smallest cost. Unfortunately, existing exact algorithms have severe scalability problems and existing approximate algorithms, though scalable, cannot guarantee near-to-optimal solutions. Motivated by this, we study the CoSKQ problem and address the above issues. Firstly, we consider the CoSKQ problem using an existing cost measurement called the *maximum sum cost*. This problem is called MaxSum-CoSKQ and is known to be NP-hard. We observe that the maximum sum cost of a set of objects is dominated by at most three objects which we call the *distance owners* of the set. Motivated by this, we propose a distance ownerdriven approach which involves two algorithms: one is an exact algorithm which runs faster than the best-known existing algorithm by several orders of magnitude and the other is an approximate algorithm which improves the best-known constant approximation factor from 2 to 1.375. Secondly, we propose a new cost measurement called the diameter cost and CoSKQ with this measurement is called Dia-CoSKQ. We prove that Dia-CoSKQ is NP-hard. With the same distance owner-driven approach, we design two algorithms for Dia-CoSKQ: one is an exact algorithm which is efficient and scalable and the other is an approximate algorithm which gives a  $\sqrt{3}$ -factor approximation. We conducted extensive experiments on real datasets which verified that the proposed exact algorithms are scalable and the proposed approximate algorithms return near-to-optimal solutions.

Worst-Case Optimized Spatial Matching. Bichromatic reverse nearest neighbor

(BRNN) queries have been studied extensively in the literature of spatial databases. Given a set P of service-providers and a set O of customers, a BRNN query is to find which customers in O are "interested" in a given service-provider in P. Recently, it has been found that this kind of queries lacks the consideration of the *capacities* of service-providers and the *demands* of customers. In order to address this issue, some spatial matching problems have been proposed, which, however, cannot be used for some real-life applications like emergency facility allocation where the maximum matching cost (or distance) should be minimized. Motivated by this, we propose a new problem called SPatial Matching for Minimizing the Maximum matching distance (SPM-MM). Then, we design two algorithms for SPM-MM, Threshold-Adapt and Swap-Chain. Threshold-Adapt is simple and easy to understand but not scalable to large datasets due to its relatively high time/space complexity. Swap-Chain, which follows a fundamentally different idea from Threshold-Adapt, runs faster than Threshold-Adapt by orders of magnitude and uses significantly less memory. We conducted extensive empirical studies which verified the efficiency and scalability of Swap-Chain.

**Direction-Preserving Trajectory Simplification.** Trajectories of moving objects are collected in many applications. Raw trajectory data is typically very large, and has to be simplified before use. Motivated by this, we introduce the notion of direction-preserving trajectory simplification, and show both analytically and empirically that it can support a broader range of applications than traditional position-preserving trajectory simplification. We present a polynomial-time algorithm for optimal direction-preserving simplification, and another approximate algorithm with a quality guarantee. Extensive experimental evaluation with real trajectory data shows the benefit of the new techniques.

The rest of this thesis is organized as follows. We introduce the collective spatial keyword query, worst-case spatial matching, and direction-preserving trajectory sim-

plification in Chapter 2, in Chapter 3 and in Chapter 4, respectively. We conclude the thesis in Chapter 5 with some future research directions.

### CHAPTER 2

# COLLECTIVE SPATIAL KEYWORD SEARCH

## 2.1 Introduction

With the proliferation of spatial-textual data such as location-based services and geotagged websites, *spatial keyword queries* have been studied extensively recently [27, 19, 67, 10]. Given a set of spatial-textual objects and a query constituted by a location and a set of keywords, a typical spatial keyword query finds the object that *best* matches the arguments in the query. One example is to find the object closest to the query location among all objects that cover all the keywords specified in the query [67].

In some applications, users' needs (expressed as keywords) are satisfied by multiple objects *collectively* instead of a *single* object [11]. For instance, a tourist wants to have site-seeing, shopping and dining which could only be satisfied by *multiple* objects, e.g., tourist attractions, shopping malls and restaurants. Another example is that a user would like to set up a project consortium of partners within a certain region that combine to offer the capabilities required for the successful execution of the whole project. Finding multiple objects collectively to satisfy users' needs can be addressed by *Collective Spatial Keyword Query* (CoSKQ) [11].

Specifically, CoSKQ is described as follows. Let  $\mathcal{O}$  be a set of objects. Each object  $o \in \mathcal{O}$  is associated with a spatial location, denoted by  $o.\lambda$ , and a set of keywords, denoted by  $o.\psi$ . Given a query q with a location  $q.\lambda$  and a set of keywords  $q.\psi$ , CoSKQ is to find a set S of objects such that S covers  $q.\psi$ , i.e.,  $q.\psi \subseteq \bigcup_{o \in S} o.\psi$ , and the cost of S, denoted by cost(S), is minimized.

There are different cost functions for cost(S). One cost function is called the  $maximum\ sum\ cost\ function$ , denoted by  $cost_{MaxSum}(S)$ , and was studied in [11]. It is the linear combination of two max components: the maximum distance between q and an object in S and the maximum distance between two objects within S. CoSKQ adopting this cost function is called MaxSum-CoSKQ. The other cost function is called the  $diameter\ cost\ function$ , denoted by  $cost_{Dia}(S)$ . It is defined to be the diameter of  $S \cup \{q\}$ . In fact, diameter-related cost functions have been commonly adopted in graph databases [2, 43, 4, 48] and spatial databases [71, 72, 73]. To the best of our knowledge, we are the first to study this cost function for CoSKQ. CoSKQ adopting this cost function is called Dia-CoSKQ.

Given a query q, an object o is said to be *relevant* (to q) if o contains at least one keyword in  $q.\psi$ . We denote by  $\mathcal{O}_q$  the set of all relevant objects to q. It is sufficient to focus on  $\mathcal{O}_q$  only for a specific query q. Given a set S of objects, S is said to be *feasible* if S covers  $q.\psi$ . Thus, the optimal solution of CoSKQ is a feasible set with the smallest cost.

Although MaxSum-CoSKQ (which is proved to be NP-hard) has been studied by Cao et al. [11], the best-known exact algorithm which we call Cao-Exact is not scalable to large datasets and the two existing approximate algorithms which we call Cao-Appro1 and Cao-Appro2 do not have a very good theoretical guarantee. Specifically, Cao-Exact is a best-first search method based on the feasible set space whose size is  $O(|\mathcal{O}_q|^{|q,\psi|})$ . Though equipped with some pruning techniques, Cao-Exact is prohibitively expensive when the dataset is large. For example, in our experiments, Cao-Exact took more than 10 days for a query containing 6 keywords on a dataset with 8M objects.

In this thesis, we propose two algorithms for MaxSum-CoSKQ, *MaxSum-Exact* and *MaxSum-Appro*. MaxSum-Exact is an exact algorithm and MaxSum-Appro is a

#### 1.375-approximate algorithm.

MaxSum-Exact is more scalable compared with the best-known algorithm, Cao-Exact. A key observation which is used by MaxSum-Exact is that the number of distinct costs of all possible feasible sets is cubic (in terms of  $|\mathcal{O}_q|$ ) although the number of all possible feasible sets is exponential (in terms of  $|q,\psi|$ ). Given a feasible set S, the maximum sum cost function of S is dominated (or determined) by at most three objects in S, namely the object with the greatest distance from q and the two objects with the greatest pairwise distance within S. We say that these three objects form the distance owner group of S. Thus, the number of distinct costs of all possible feasible sets is bounded by the total number of all possible distance owner groups (which is bounded by  $O(|\mathcal{O}_q|^3)$ ). Motivated by this, we propose a distance-owner driven approach called MaxSum-Exact for MaxSum-CoSK. MaxSum-Exact is a search algorithm based on the search space containing all possible distance owner groups. Besides, it incorporates some search strategies which can prune the search space effectively. Usually, one distance owner group corresponds to many feasible sets. This is verified by our experiments where MaxSum-Exact ran faster than Cao-Exact by 1-3 orders of magnitude.

MaxSum-Appro, the proposed approximate algorithm, improves the best-known constant approximation factor from 2 to 1.375 without incurring a higher worst-case time complexity.

Furthermore, we consider Dia-CoSKQ which has not been studied in the literature. In this thesis, we prove that Dia-CoSKQ is NP-hard. We also adapt Cao-Exact, Cao-Appro1 and Cao-Appro2 for Dia-CoSKQ. However, these adapted algorithms suffer from the same drawbacks in MaxSum-CoSKQ.

Motivated by this, we propose two algorithms, namely *Dia-Exact* and *Dia-Appro*. Dia-Exact is an exact algorithm which is also a search algorithm based on the search space containing all possible *distance owner groups* and thus it is scalable to large

datasets. Dia-Appro gives a  $\sqrt{3}$ -factor approximation for Dia-CoSKQ.

We summarize our main contributions as follows.

- Firstly, for MaxSum-CoSKQ, we design two algorithms, MaxSum-Exact and MaxSum-Appro. MaxSum-Exact is more scalable than the best-known exact algorithm, Cao-Exact. MaxSum-Appro improves the best-known constant approximation factor from 2 to 1.375 without incurring a higher worst-case time complexity.
- Secondly, for Dia-CoSKQ, which is new, we prove its NP-hardness and develop two algorithms, Dia-Exact and Dia-Appro. Dia-Exact significantly outperforms the adaptation of Cao-Exact, and Dia-Appro gives a √3-factor approximation.
- Thirdly, we conducted extensive experiments on both real and synthetic datasets, which verified our theoretical results and the efficiency of our algorithms.

The rest of this chapter is organized as follows. Section 2.2 gives the definition of the CoSKQ problem and its existing solutions. Section 2.3 and Section 2.4 study MaxSum-CoSKQ and Dia-CoSKQ, respectively. Section 2.5 gives the empirical study and Section 2.6 reviews the related work. Section 2.7 concludes this chapter.

# 2.2 Background

### 2.2.1 Problem Definition

Let  $\mathcal{O}$  be a set of objects. Each object  $o \in \mathcal{O}$  is associated with a location denoted by  $o.\lambda$  and a set of keywords denoted by  $o.\psi$ . Given two objects o and o', we denote by d(o,o') the Euclidean distance between  $o.\lambda$  and  $o'.\lambda$ . Given a query q which consists of a location  $q.\lambda$  and a set of keywords  $q.\psi$ , we denote by  $\mathcal{O}_q$  the set of **relevant** objects each of which contains at least one keyword in  $q.\psi$ , and say that a set of objects is

**feasible** if it covers  $q.\psi$ . Besides, we introduce a fictitious object  $o_q$  in  $\mathcal{O}$  with  $o_q.\lambda = q.\lambda$  and  $o_q.\psi = \emptyset$ . For simplicity, we shall also refer to object  $o_q$  as q.

**Problem Definition [11].** Given a query  $q = (q.\lambda, q.\psi)$ , the *Collective Spatial Keyword Query* (CoSKQ) problem is to find a set S of objects in  $\mathcal{O}$  such that S covers  $q.\psi$  and the *cost* of S is minimized.

In this thesis, we consider two cost functions, the *maximum sum cost* and the *diameter cost*.

Given a set S of objects, the maximum sum cost of S, denoted by  $cost_{MaxSum}(S)$ , is equal to the linear combination of the maximum distance between q and an object in S and the maximum distance between two objects in S. That is,

$$cost_{MaxSum}(S) = \alpha \cdot \max_{o \in S} d(o, q) + (1 - \alpha) \cdot \max_{o_1, o_2 \in S} d(o_1, o_2)$$
 (2.1)

where  $\alpha \in [0,1]$  is a user parameter. Same as [11], for ease of exposition, we consider the case where  $\alpha = 0.5$  only. In this case, we can safely assume that

$$cost_{MaxSum}(S) = \max_{o \in S} d(o, q) + \max_{o_1, o_2 \in S} d(o_1, o_2)$$
 (2.2)

In fact, the applicability of all of our algorithms does not rely on the setting of  $\alpha$ . The only part that is affected is the approximation factor of our approximate algorithm which is *bounded* by  $(2-\sqrt{2}/2\cdot\alpha)$  (e.g., when  $\alpha=0.5$ , the approximation factor of our approximate algorithm is 1.375 which is bounded by  $(2-\sqrt{2}/2\cdot\alpha)\approx 1.65$ ). More details in Section A.1 of the appendix. The CoSKQ problem using this cost is called *MaxSum-CoSKQ*.

As could be noticed, parameter  $\alpha$  in the maximum sum cost function is used to balance the two max components, namely  $\max_{o \in S} d(o,q)$  and  $\max_{o_1,o_2 \in S} d(o_1,o_2)$ . Sometimes, however, people may not have a concrete idea of how to specify  $\alpha$ . To

ease this situation, we define an alternative cost function called *diameter cost* on a set S of objects, denoted by  $cost_{Dia}(S)$ , which is defined to be the larger of these two max components. That is,

$$cost_{Dia}(S) = \max_{o_1, o_2 \in S \cup \{o_q\}} d(o_1, o_2)$$
(2.3)

The CoSKQ problem using this cost is called *Dia-CoSKQ*.

**Intractability.** It has been proved in [11] that MaxSum-CoSKQ is NP-hard. In this thesis, we prove that Dia-CoSKQ is also NP-hard.

**Proof.** We first give the decision problem of Dia-CoSKQ. Let  $\mathcal{O}$  be a set of spatial objects each of which is associated with a set of keywords. Given a number C and a query q consisting of a location  $q.\lambda$  and a set of keywords  $q.\psi$ , the problem is to determine whether there exists a set S of objects such that S covers  $q.\psi$  and  $cost_{Dia}(S) < C$ . For simplicity, we denote this decision problem by Dia-CoSKQ.

We then utilize a well-known NP-C problem 3-SAT for proving that Dia-CoSKQ is NP-hard. 3-SAT is described as follows. Let U be a set of literals (binary variables)  $\{e_1, \overline{e_1}, ..., e_n, \overline{e_n}\}$ . Note that  $\overline{e_i}$  corresponds to the negation of  $e_i$ . Given an expression  $E = C_1 \wedge C_2 \wedge ... \wedge C_m$  where  $C_i$  has its from of  $x_i \vee y_i \vee z_i$  and  $x_i, y_i, z_i \in U$  for  $1 \leq i \leq m$ , it determines whether there exists a truth assignment for  $e_i$  for  $1 \leq i \leq n$  such that E is true.

Given a 3-SAT problem instance, we construct a Dia-CoSKQ problem instance as follows. We arbitrarily select a location as  $q.\lambda$  and for each clause  $C_i$  in the expression E, we include  $C_i$  in  $q.\psi$  as a keyword. Thus, q contains m keywords. To construct  $\mathcal{O}$ , we consider the circle Cir with its center at  $q.\lambda$  and its radius equal to 1. For each pair of literals  $e_i$  and  $\overline{e_i}$  in U, we create two objects  $o_i$  and  $o'_i$  on the boundary of Cir

such that  $d(o_i, o'_i) = 2$ , i.e., the line segment between  $o_i$  and  $o'_i$  is a diameter of Cir. We guarantee that no two objects in  $\mathcal{O}$  share the same location. Besides, we set  $o_i.\lambda$   $(o'_i.\lambda)$  to be the set of clauses that contain  $e_i$  ( $\overline{e_i}$ ). We set C to be 2. Clearly, the above construction process could be finished in polynomial time.

We proceed to show that the above constructed Dia-CoSKQ problem instance is equivalent to its corresponding 3-SAT problem instance. Assume that the answer to 3-SAT is "yes", i.e., there exists a truth assignment A for the literals in U such that E is true. Then, we construct a set S of objects in  $\mathcal{O}$  as follows. For each positive literal  $e_i \in U$ , we include  $o_i$  in S if  $e_i$  is true in A. For each negative literal  $\overline{e_i} \in U$ , we include  $o'_i$  in S if  $e_i$  is false in A. Clearly,  $o_i$  and  $o'_i$  cannot be included in S simultaneously. Consider S. First, S covers  $q.\psi$  which could be verified by contradiction. Assume that there exists a keyword  $C_i$  in  $q.\psi$  which is not covered by S. Since E is true,  $C_i$  must also be true. Consequently,  $C_i$  contains at least one literal which is true. Case 1: this literal is a positive literal  $e_i$ . In this case,  $o_i$  is included in S and thus S covers  $C_i$ , which leads to a contradiction. Case 2: this literal is a negative literal  $\overline{e_i}$ . In this case,  $e_i$  is false and thus  $o'_i$  is included in S. As a result, S covers  $C_i$ , which, again, leads to a contradiction. Second, we know  $cost_{Dia}(S) < C = 2$ . This is because  $\max_{o \in S} d(o) = 1$  and  $\max_{o_1,o_2 \in S} d(o_1,o_2) < 2$  (there exist no pairs of two objects such that the segment between them forms a diameter).

Consider the other direction. Assume that the answer to Dia-CoSKQ is "yes", i.e., there exist a set S of objects such that S covers  $q.\psi$  and  $cost_{Dia}(S) < C$ . Then, we construct a truth assignment A for the literals in U as follows. For each object  $o_i \in S$ , we set  $e_i$  to be true and consequently  $\overline{e_i}$  is false. For each object  $o_i' \in S$ , we set  $e_i$  to be false and consequently  $\overline{e_i}$  is true. For the remaining literals to which no truth values have been assigned, we set their truth values arbitrarily with the constraint that  $e_i$  and  $\overline{e_i}$  have different truth values. First, we show that E is a *valid* assignment, i.e.,

there exist no pairs of two literals  $e_i$  and  $\overline{e_i}$  such that they have a common truth value. This could be verified by contradiction. Assume that there exist two literals  $e_i$  and  $\overline{e_i}$  such that  $e_i$  and  $\overline{e_i}$  have the same truth value. It follows that  $o_i$  and  $o_i'$  co-exist in S which contradicts the assumption that  $cost_{Dia}(S) < C = 2$ . Second, we show that E is true with the truth assignment A. Again, this could be verified by contradiction. Assume that E is false. Then, there exists a clause  $C_i$  which is false. It follows that each positive literal  $e_i$  in  $C_i$  (if any) is false and thus  $o_i$  is not included in S. Each negative literal  $\overline{e_i}$  in  $C_i$  (if any) is false and thus  $o_i'$  is not included in S. As a result, S does not cover  $C_i$  which contradicts that S covers  $q.\psi$ . Therefore, E is true and thus the answer to the 3-SAT problem is "yes".

Therefore, we know that Dia-CoSKQ is NP-hard. ■

### 2.2.2 Existing Solutions for MaxSum-CoSKQ

Cao et el. [11] proposed one exact algorithm, Cao-Exact, and two approximate algorithms, Cao-Appro1 and Cao-Appro2, for MaxSum-CoSKQ.

Cao-Exact. Cao-Exact is a best-first search method using an index called IR-tree [19]. An IR-tree is an R-tree in which each node is augmented with an  $Inverted\ File\ (IF)$ . Consider a leaf node N. For each keyword t, we construct an  $inverted\ list$  which is a list of all objects in node N containing t. All inverted lists in this leaf node N form the IF of N. Consider a non-leaf node N'. For each keyword t, we construct an  $inverted\ list$  which is a list of all child nodes in N' covering t. Given a keyword t, a node N'' is said to cover t if there exists an object in the subtree rooted at N'' containing t. All inverted lists in this non-leaf node N' form the IF of N'.

Cao-Exact is basically an exhaustive search on the object space with some pruning strategies in the IR-tree. The worst-case time complexity of Cao-Exact is  $O(|\mathcal{O}|^{|q.\psi|})$ , which corresponds to the size of the set containing all possible feasible sets.

**Cao-Appro1.** Cao-Appro1 gives a 3-factor approximation for MaxSum-CoSKQ. Specifically, Cao-Appro1 finds for each  $t \in q.\psi$ , q's nearest neighbor (NN) in  $\mathcal{O}$  containing t and returns the set containing all these NNs as the approximate solution. Since Cao-Appro1 issues NN queries at most  $|q.\psi|$  times and each NN query takes  $O(\log |\mathcal{O}|)$  time [13, 22, 52], the time complexity of Cao-Appro1 is  $O(|q.\psi| \cdot \log |\mathcal{O}|)$ . Cao-Appro2. Cao-Appro2 gives a 2-factor approximation for MaxSum-CoSKQ. Specifically, Cao-Appro2 enhances Cao-Appro1 as follows. First, Cao-Appro2 invokes Cao-Appro1 and obtains an approximate solution denoted by  $S_1$ . Let  $o_f$  be the farthest object from q in  $S_1$  and  $t_f$  be a keyword contained by  $o_f$  but not contained by any other closer object from q in  $\mathcal{O}$ . Then, for each object o in  $\mathcal{O}$  containing  $t_f$ , it finds for each keyword t in  $q.\psi$ , o's nearest object that contains t in  $\mathcal{O}$  and obtains a corresponding approximate solution containing all these NNs. Among all these approximate solutions as well as  $S_1$ , it returns the one with the smallest cost. Thus, the approximate solution returned by Cao-Appro2 is no worse than that returned by Cao-Appro1. Since there are at most  $|\mathcal{O}_q|$  objects containing  $t_f$  and the cost for each such object is simply  $O(|q.\psi| \cdot \log |\mathcal{O}|)$ , the worst-case time complexity of Cao-Appro2 is

# 2.3 Algorithms for MaxSum-CoSKQ

 $O(|\mathcal{O}_q| \cdot |q.\psi| \cdot \log |\mathcal{O}|).$ 

In this section, we propose two algorithms, MaxSum-Exact (Section 2.3.1) and MaxSum-Appro (Section 2.3.2), for MaxSum-CoSKQ. For clarity, we simply write  $cost_{MaxSum}(\cdot)$  as  $cost(\cdot)$  if the context of the cost function is clear.

Given a query q and a non-negative real number r, we denote the *circle* or the *disk* centered at  $q.\lambda$  with radius r by D(q,r). Given a disk D, we denote the radius of D by radius(D). Given a query q, a disk centered at  $q.\lambda$  is called a q-disk. Given a q-disk D and an object o in D, o is said to be the *boundary object* of D if there does not exist

other objects o' in D such that d(o',q) > d(o,q). Note that in some cases, a boundary object of a disk is along the boundary of a disk and in some other cases, it is inside the disk without touching the boundary of the disk.

### 2.3.1 Finding Optimal Solution

In this section, we propose an exact algorithm called *MaxSum-Exact*. The key to the efficiency of MaxSum-Exact is based on the splitting property of the maximum sum cost function.

#### **Splitting Property**

Let S' be a feasible set. The maximum sum cost of S' can be split into two parts, namely the *query distance cost* which is  $\max_{o \in S'} d(o, q)$  and the *pairwise distance cost* which is  $\max_{o_1,o_2 \in S'} d(o_1,o_2)$ . We define the **query distance owner** of S' to be o where  $o = \arg \max_{o \in S'} d(o,q)$ . We also define the **pairwise distance owners** of S' to be  $o_1$  and  $o_2$  where  $(o_1,o_2) = \arg \max_{(o'_1,o'_2) \in S' \times S'} d(o'_1,o'_2)$ .

Consider Figure 2.1 containing a query location q and 5 objects, namely  $o_1, o_2, o_3, o_4$  and  $o_5$ . The set of keywords associated with each object can be found in the figure. Suppose that  $q.\psi = \{t_1, t_2, t_3\}$ . We know that a set  $S' = \{o_1, o_2, o_3\}$  is feasible. The query distance owner of S' is  $o_1$  and the pairwise distance owners of S' are  $o_2$  and  $o_3$ .

According to the above splitting property, the cost of a set S' can be dominated (or determined) by exactly three objects in S', namely the query distance owner of S' (i.e., o) and the two pairwise distance owners of S' (i.e.,  $o_1$  and  $o_2$ ). In other words, we can simply write the cost of S' as follows.

$$cost(S') = d(o,q) + d(o_1, o_2)$$

where o is the distance owner of S', and  $o_1$  and  $o_2$  are the two pairwise distance owners of S'. We say that o,  $o_1$  and  $o_2$  forms a **distance owner group**. Any feasible set with its query distance owner as o and its pairwise distance owners as  $o_1$  and  $o_2$  is said to be  $(o, o_1, o_2)$ -owner consistent. Note that each feasible set that is  $(o, o_1, o_2)$ -owner consistent has the same cost equal to  $d(o, q) + d(o_1, o_2)$ .

#### **Distance Owner-Driven Approach**

Based on the splitting property, we propose a distance owner-driven approach as follows. This approach maintains a variable S storing the best feasible set found so far. Initially, S is set to a feasible set (We will describe how we find this feasible set later). Then, it has four major steps.

- Step 1 (Query Distance Owner Finding): Select one object o in  $\mathcal{O}_q$  to take the role of the query distance owner of a set S' to be found.
- Step 2 (Pairwise Distance Owner Finding): Select two objects,  $o_1$  and  $o_2$ , in  $\mathcal{O}_q$  to take the roles of the pairwise distance owners of the set S' (to be found). Note that  $o, o_1$  and  $o_2$  form a distance owner group.
- Step 3 (Sub-Optimal Feasible Set Finding): Find the set S' which is  $(o, o_1, o_2)$ owner consistent (if any), and update S with S' if cost(S') < cost(S).
- Step 4 (Iterative Step): Repeat Step 1 and Step 2 which find another distance owner group, and continue with Step 3 until all distance owner groups are traversed.

The above approach gives a search strategy based on the set of all possible distance owner groups. However, a straightforward implementation of this approach would enumerate all  $|\mathcal{O}_q|^3$  distance owner groups, which is prohibitively expensive in practice. Thus, we need a careful design in order to prune the search space effectively. In

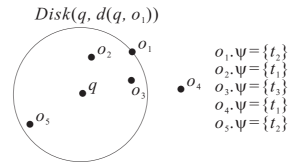


Figure 2.1: An example (for CoSKQ)

the following, we elaborate the pruning features enjoyed by this distance owner-driven approach, which cannot be found in the best-known algorithm, Cao-Exact.

Firstly, some objects in  $\mathcal{O}_q$  need not be considered in Step 2 after we select an object in Step 1. To illustrate this, consider Figure 2.1. Suppose that we pick  $o_1$  as the query distance owner in Step 1. We do not need to consider  $o_4$  as objects in Step 2. This is because  $d(o_4, q)$  is larger than  $d(o_1, q)$ , which violates the property that  $o_1$  takes the role of the query distance owner of the set S' to be found if S' contains  $o_1$  and  $o_4$ . We formalize this pruning feature as follows.

**Property 1 (Pruning)** Let S' be a feasible set. If o is the query distance owner of S', then the two pairwise distance owners of S' are inside D(q, d(o, q)).

**Proof.** Any object  $o' \in S'$  has  $d(o', q) \leq d(o, q)$  and thus o' is inside D(q, d(o, q)).

Secondly, most of the objects in  $\mathcal{O}_q$  need not be considered to form a set S' to be found in Step 3. To illustrate this, consider Figure 2.1 again. Suppose that we pick  $o_1$  as the query distance owner in Step 1, and  $o_2$  and  $o_3$  as the pairwise distance owners in Step 2. Similarly, we still do not need to consider  $o_4$  as one of the objects to form the set S' since including  $o_4$  violates the query distance owner property. Besides, we do not need to consider  $o_5$  to form the set S' to be found. This is because  $d(o_2, o_5) > d(o_2, o_3)$  which violates the property that  $o_2$  and  $o_3$  take the roles of the pairwise distance owners. Similarly, we formalize this pruning feature as follows.

**Property 2 (Pruning)** Let S' be a feasible set. If o is the query distance owner of S', and  $o_1$  and  $o_2$  are two pairwise distance owners of S', then all objects in S' are inside  $\mathcal{R}$  where  $\mathcal{R} = D(q, d(o, q)) \cap D(o_1, d(o_1, o_2)) \cap D(o_2, d(o_1, o_2))$ .

**Proof.** For each  $o' \in S'$ , we have  $d(o',q) \leq d(o,q)$  which implies that o' is inside D(q,d(o,q)). For each  $o' \in S'$ , we have  $d(o',o_1) \leq d(o_1,o_2)$  which implies that o' is inside  $D(o_1,d(o_1,o_2))$ , and  $d(o',o_2) \leq d(o_1,o_2)$  which implies that o' is inside  $D(o_2,d(o_1,o_2))$ .

The above pruning features look promising for improving the efficiency of the proposed approach. Moreover, since objects near to q usually form the optimal set, we propose to consider the objects in Step 1 iteratively, taking the role of the query distance owner of the set to be found, in ascending order of their distances to q in order to further improve the efficiency of the proposed approach.

Usually, the NN of q in  $\mathcal{O}_q$  is not the query distance owner of the set S' to be found. In Figure 2.1, consider the query q with its keyword set to be  $\{t_1, t_2, t_3\}$ . The NN of q is  $o_2$ . Suppose that  $o_2$  is the query distance owner of S'. According to Property 2, all objects in S' fall in  $D(q, d(o_2, q))$  and they together cover  $q.\psi$ . But, in the figure, no object in  $D(q, d(o_2, q))$  contains  $t_2$ , which implies that we cannot find a feasible set S' with  $o_2$  as its query distance owner.

Based on this observation, we propose to find the *closest possible query distance* owner, say o, of the set S' to be found such that there exists a feasible set in the q-disk D(q, d(o, q)). In addition, we do not want to pick any object which is far away from q. Thus, we also propose to find the *farthest possible query distance owner* of S' to be found that we need to consider.

#### Closest/Farthest Possible Query Dist. Owner

The following two lemmas show how to find the closest and farthest possible query distance owners.

Before we present the first lemma about the closest possible query distance owner, we introduce some notations. Given a query q and a keyword t, the t-keyword nearest neighbor of q, denoted by NN(q,t), is defined to be the NN of q containing keyword t. We have a similar definition on NN(o,t) for an object o. We define the nearest neighbor set of q, denoted by N(q), to be the set containing q's t-keyword nearest neighbor for each  $t \in q.\psi$ , i.e., N(q) is  $\cup_{t \in q.\psi} NN(q,t)$ . Note that N(q) is a feasible set.

**Lemma 2.3.1** (Closest Poss. Query Dist. Owner) Let  $r_{min} = \max_{o \in N(q)} d(o, q)$ .

There exists a feasible set in a q-disk D if and only if  $radius(D) \ge r_{min}$ .

**Proof.** The proof for the "if" part is trivial since for any q-disk D with  $radius(D) \geq r_{min}$ , N(q) is a feasible set in D. We prove the "only if" part by contradiction. Assume  $radius(D) < r_{min}$  and there exists a feasible set S in D. Let  $o_f$  be the farthest object from q in N(q), i.e.,  $r_{min} = d(q, o_f)$ . There exists a keyword  $t_f \in o_f . \psi \cap q . \psi$  such that  $t_f$  is not contained by any object that is closer to q than  $o_f$  since otherwise  $o_f \notin N(q)$ . Since S is feasible, there exists an object  $o \in S$  that contains keyword  $t_f$ . As a result, we have  $d(o,q) \leq radius(D) < r_{min} = d(q,o_f)$ , which leads to a contradiction.  $\blacksquare$ 

The above lemma suggests that there is no feasible set in a q-disk D if  $radius(D) < r_{min}$ . Thus, the disk with its radius equal to  $r_{min}$  is the "smallest" disk we need to consider. The boundary object of this disk is the closest possible query distance owner. Note that this object is along the boundary of this disk.

The following lemma gives the "largest" disk we need to consider. Besides, the boundary object of this disk corresponds to the farthest possible query distance owner.

Note that this object might or might not be along the boundary of this disk.

**Lemma 2.3.2 (Farthest Poss. Query Dist. Owner)** Let S be a feasible set and  $r_{max} = cost(S)$ . Let D be a q-disk with  $radius(D) > r_{max}$ . Then, for any feasible set S' containing at least one object outside D, cost(S') > cost(S).

**Proof.** 
$$cost(S') \ge max_{o \in S'}d(o,q) > radius(D) > r_{max} = cost(S)$$
.

The above lemma suggests that when we have known a feasible set S, there is no need to consider the objects outside  $D(q, r_{max})$  where  $r_{max} = cost(S)$ .

The above two lemmas suggest the "smallest" disk and the "largest" disk we need to consider. Specifically, the object o which takes the role of the query distance owner of S' to be found must be in the ring which is roughly equal to the "largest" disk minus the "smallest" disk. Let S be a feasible set. Let  $r_{min} = \max_{o \in N(q)} d(o, q)$  and  $r_{max} = cost(S)$ . We define the **ring** for S, denoted by R(S), to be  $D(q, r_{max}) - D(q, r_{min} - \delta)$ , where  $\delta$  is a very small positive real number near to 0.

**Lemma 2.3.3** (Ring Candidate) Let S be a feasible set and  $S_o$  be the optimal set for the MaxSum-CoSKQ problem. The query distance owner of  $S_o$  is inside R(S).

**Proof.** Let o be the query distance owner of  $S_o$ . First, according to Lemma 2.3.2, o cannot be outside  $D(q, r_{max})$  since otherwise  $cost(S_o) > cost(S)$  which leads to a contradiction. Second, according to Lemma 2.3.1, there exist no feasible sets in  $D(q, r_{min} - \delta)$ . Thus, o is not inside  $D(q, r_{min} - \delta)$  since otherwise  $S_o$  which is feasible is inside  $D(q, r_{min} - \delta)$  which also leads to a contradiction. Therefore, o is inside R(S).

It is easy to verify that the region occupied by R(S) becomes smaller when cost(S) is smaller since the radius of the outer disk of R(S) is equal to cost(S).

#### The MaxSum-Exact Algorithm

Based on the discussion in the previous subsection, we design MaxSum-Exact as shown in Algorithm 1. Specifically, we maintain S for storing the best-known solution found so far, which is initialized to N(q). Then, we perform an iterative process as follows. Consider an iteration. We want to check whether there exists a relevant object in R(S)that has not been processed. If yes, we pick the nearest relevant object o from R(S)that has not been processed to take the role of the query distance owner of the set S' to be found (Step 1). This object is said to be the *query distance owner* for this iteration. We process it as follows. Firstly, we form the q-disk D with its radius equal to d(o, q)and find a set P of all pairs  $(o_1, o_2)$  where  $o_1$  and  $o_2$  are in D for taking the roles of the pairwise distance owners (Step 2). Secondly, for each pair  $(o_1, o_2)$  in P which is processed in ascending order of  $d(o_1, o_2)$ , we check whether there exists a feasible set S' which is  $(o, o_1, o_2)$ -owner consistent. Case 1: yes. We do the following. Firstly, if cost(S') < cost(S), then we update S by S'. Secondly, we terminate to search the remaining pairs in P since the cost of a final set whose pairwise distance owners corresponds to one of the remaining pairs must be at least the cost of the current set S' whose pairwise distance owners are  $(o_1, o_2)$ , the current processed pair. Case 2: no. We continue to consider the next pair in P until Case 1 is reached or all the pairs in P have been processed. We continue the above iteration with the next relevant object from R(S) that has not been processed until all objects in R(S) have been processed (Step 4).

We verify the correctness of MaxSum-Exact via Theorem 2.3.1.

Theorem 2.3.1 MaxSum-Exact returns a feasible set with the smallest cost for MaxSum-CoSKQ. □

**Proof.** Let  $S_o$  be one of the feasible sets with the smallest cost for MaxSum-CoSKQ.

### Algorithm 1 Algorithm MaxSum-Exact

```
Require: query q and a set \mathcal{O} of objects
 1: S \leftarrow N(q)
 2: while there is an "un-processed" relevant object o in R(S) do
      // Step 1 (Query Distance Owner Finding)
      o \leftarrow the nearest "un-processed" relevant object in R(S)
 4:
 5:
      // Step 2 (Pairwise Distance Owner Finding)
       D \leftarrow the q-disk with its radius equal to d(o, q)
 6:
       P \leftarrow a set of all pairs (o_1, o_2) where o_1 and o_2 are in D
 7:
       // Step 3 (Sub-optimal Feasible Set Finding)
 8:
       for each (o_1, o_2) \in P in ascending order of d(o_1, o_2) do
 9:
         if there exists a feasible set S' in D which is (o, o_1, o_2)-owner consistent then
10:
            if cost(S') < cost(S) then
11:
               S \leftarrow S'; break
12:
       // Step 4 (Iterative Process)
13:
       mark o as "processed"
14:
15: return S
```

Suppose that o is the query distance owner of  $S_o$ , and  $o_1$  and  $o_2$  are two pairwise distance owners of  $S_o$ . According to Lemma 2.3.3, o is inside R(S), where S is the solution maintained in MaxSum-Exact. Thus, o must have been processed in MaxSum-Exact (Step 1). When o is processed, pair  $(o_1, o_2)$  is included in P (Step 2) since  $o_1$  and  $o_2$  are inside D(q, d(o, q)) (Property 1). As a result, any feasible set which is  $(o, o_1, o_2)$ -owner consistent is retrieved (Step 3) and used to update S (there must exist some since  $S_o$  is  $(o, o_1, o_2)$ -owner consistent). The resulting S will not be updated anymore since it has the same cost as  $cost(S_o)$  which is the smallest, and thus S is the final output.  $\blacksquare$ 

Algorithm 1 looks straightforward but how to execute this algorithm *efficiently* needs more careful design. We propose two computation strategies in the algorithm, namely the *self-iteration computation strategy* and the *cross-iteration computation strategy*, to execute this algorithm efficiently. The self-iteration computation strategy is to speed up the operations within an iteration and the cross-iteration computation strategy is to speed up the operations across different iterations.

**Self-Iteration Computation Strategy.** Consider an iteration in the algorithm whose query distance owner is o. Step 1 (lines 3-4) is straightforward. In Step 2 (lines 5-7),

there is a step of finding a set P of all pairs  $(o_1, o_2)$  where  $o_1$  and  $o_2$  are in D. There is no need to keep all pairs  $(o_1, o_2)$  in P and some pairs can be pruned. The following two lemmas give some hints for pruning. The first lemma (Lemma 2.3.4) is based on the triangle inequality and the second lemma (Lemma 2.3.5) is based on the best-known set S found so far.

**Lemma 2.3.4 (Triangle Inequality)** Let S' be a feasible solution whose query distance owner is o, and pairwise distance owners are  $o_1$  and  $o_2$ . Then,  $d(o_1, o_2) \ge d(o, q) - \min\{d(o_1, q), d(o_2, q)\}$ .

**Proof.** Note that  $d(o_1, o_2) \ge d(o_1, o)$  and  $d(o_1, o_2) \ge d(o_2, o)$ . By the triangle inequality, we know  $d(o_1, o) \ge d(o, q) - d(o_1, q)$  and  $d(o_2, o) \ge d(o, q) - d(o_2, q)$ . Thus, we have  $d(o_1, o_2) \ge d(o, q) - \min\{d(o_1, q), d(o_2, q)\}$ .

The above lemma suggests that the pair  $(o_1, o_2)$  in P can be pruned if  $d(o_1, o_2) < d(o, q) - \min\{d(o_1, q), d(o_2, q)\}$ . Let  $d_{min} = d(o, q) - \min\{d(o_1, q), d(o_2, q)\}$ . Thus,  $d_{min}$  corresponds to the smallest distance threshold for a pair  $(o_1, o_2)$ .

**Lemma 2.3.5 (Best Known Set)** Let S' be a feasible solution whose query distance owner is o and pairwise distance owners are  $o_1$  and  $o_2$ . Let S be another feasible solution.  $cost(S') \leq cost(S)$  if and only if  $d(o_1, o_2) \leq cost(S) - d(o, q)$ .

**Proof.**  $cost(S') \leq cost(S)$  deduces  $d(o,q) + d(o_1,o_2) \leq cost(S)$  which is exactly  $d(o,q) \leq cost(S) - d(o_1,o_2)$ .

Let S be the feasible set found so far in the algorithm. The above lemma suggests that the pair  $(o_1, o_2)$  in P can be pruned if  $d(o_1, o_2) > cost(S) - d(o, q)$ . Let  $d_{max} = cost(S) - d(o, q)$ . Thus,  $d_{max}$  is the largest distance threshold for a pair  $(o_1, o_2)$ .

According to Lemma 2.3.4 and Lemma 2.3.5, we only need to maintain those pairs with their distances between  $d_{min}$  and  $d_{max}$  in P.

Consider Step 3 (lines 8-12). Here, we need to process each pair  $(o_1, o_2)$  in P. The most time-consuming operation is to check whether there exists a feasible set S' which is  $(o, o_1, o_2)$ -owner consistent. Algorithm 2 presents an algorithm for this task. If it succeeds, it outputs S'; otherwise, it outputs  $\emptyset$ . First, it checks whether  $d(o_1, o_2) < \max\{d(o_1, o), d(o_2, o)\}$ . If yes, we conclude that there exist no feasible set that is  $(o, o_1, o_2)$ -owner consistent since it violates the condition that  $o_1$  and  $o_2$  are the pairwise distance owners (i.e.,  $d(o_1, o_2) \ge \max\{d(o_1, o), d(o_2, o)\}$ ). If no, it initializes S' to be  $\{o, o_1, o_2\}$ . It also maintains a variable  $\psi$ , denoting the set of keywords not covered by S' yet, which is initialized as  $q.\psi - (o.\psi \cup o_1.\psi \cup o_2.\psi)$ . If  $\psi = \emptyset$ , it returns S' immediately. Otherwise, it proceeds to augment S' with some other objects. According to Property 2, we can safely focus on the region  $\mathcal{R} = D(o, d(o, q)) \cap$  $D(o_1, d(o_1, o_2)) \cap D(o_2, d(o_1, o_2))$ . Therefore, it retrieves the set  $\mathcal{O}'$  of all relevant objects in  $\mathcal{R}$ . If  $\mathcal{O}'$  does not cover  $\psi$ , it returns  $\emptyset$ . Otherwise, it enumerates each possible subset S'' of  $\mathcal{O}'$  that covers  $\psi$  (by utilizing the *inverted lists* maintained for each keyword in  $\psi$ ), augment S' by S'' (thus S' becomes feasible) and checks whether S' is  $(o, o_1, o_2)$ -owner consistent which is equivalent to checking whether  $o_1$  and  $o_2$  are still the pairwise distance owners of S'. If yes, it outputs S'. Otherwise, it restores S' and checks the next subset of  $\mathcal{O}'$ . When all subsets of  $\mathcal{O}'$  that cover  $\psi$  have been traversed and still no feasible set S' which is  $(o, o_1, o_2)$ -owner consistent has been found, it returns  $\emptyset$ .

**Cross-Iteration Computation Strategy.** We reuse the information computed in the previous iterations for the current iteration.

Consider an iteration where the query distance owner for this iteration is o. With respect to o, we create a q-disk D and also construct set P (line 7 in Algorithm 1). Consider the next iteration where the query distance owner for this iteration is o'. Although we can construct set P' with respect to o' from scratch by applying the procedure of

**Algorithm 2** Algorithm for checking whether there exists a feasible set S' which is  $(o, o_1, o_2)$ -owner consistent

```
Require: three objects o, o_1 and o_2

Ensure: a feasible set which is (o, o_1, o_2)-owner consistent if any and \emptyset otherwise

1: if d(o_1, o_2) < \max\{d(o_1, o), d(o_2, o)\} then return \emptyset

2: S' \leftarrow \{o, o_1, o_2\}

3: \psi \leftarrow q.\psi - (o.\psi \cup o_1.\psi \cup o_2.\psi)

4: if \psi = \emptyset then return S'

5: \mathcal{R} \leftarrow D(q, d(o, q)) \cap D(o_1, d(o_1, o_2)) \cap D(o_2, d(o_1, o_2))

6: \mathcal{O}' \leftarrow a set of all relevant objects in \mathcal{R}

7: if \mathcal{O}' does not cover \psi then return \emptyset

8: for each subset S'' of \mathcal{O}' that covers \psi do

9: S' \leftarrow S' \cup S''

10: if S' is (o, o_1, o_2)-owner consistent then return S'

11: S' \leftarrow S' - S''

12: return \emptyset
```

generating set P, a much better approach is to construct set P' by using the current content of P because  $P \subseteq P'$ . Specifically, when we consider the next iteration, we first construct another set Q to be the set of additional pairs in P' compared with P (i.e.,  $Q = \{(o'', o') | o'' \in D(q, d(q, o))\}$ ) and then set P' to be  $P \cup Q$ . Note that  $P \cap Q = \emptyset$ .

The pruning in P mentioned in the self-iteration computation strategy is still valid even when we construct P' in the above way. Specifically, the pairs pruned previously in P still do not need to be considered in P' at the next iteration. This is because  $d_{min}$  is monotonically increasing and  $d_{max}$  is monotonically decreasing with more iterations. To illustrate, consider a pair  $(o_1,o_2)$  in P at the previous iteration. Note that  $d_{min} = d(o,q) - \min\{d(o_1,q),d(o_2,q)\}$  and  $d_{max} = cost(S) - d(o,q)$ . At the next iteration, o will become o', which is at least as far as o from q (i.e.,  $d(o',q) \ge d(o,q)$ ). Thus, at the next iteration,  $d_{min}$  will remain the same or will increase. In addition, the cost of the solution S maintained at the next iteration is at most the cost of that maintained at the previous iteration. Thus, at the next iteration,  $d_{max}$  will remain the same or will decrease.

### **Implementation and Time Complexity**

We adopt the IR-tree built on  $\mathcal{O}$  to support both the NN query (line 1 of Algorithm 1) and the range query (line 7 of Algorithm 1 and line 6 of Algorithm 2). For the NN query, we adopt the best-first search method [34] and for the range query, we perform a simple breadth-first traversal with the constraint of the range. Besides, given a query q, since we only focus on the set of relevant objects, when performing NN queries and range queries, we can utilize the IF information maintained in the IR-tree for pruning.

Since the pairs in P are processed in ascending order of their distances and P is maintained dynamically (because of the Cross-Iteration Computation Strategy), we adopt a *binary search tree* for maintaining P, which allows efficient sorting and update.

Let  $n_1$  be the number of iterations (lines 2-14) in MaxSum-Exact (Algorithm 1). Note that  $n_1 << |\mathcal{O}_q|$  since  $n_1$  corresponds to the number of relevant objects we process in R(S) and the area occupied by R(S) is typically small. Let |P| be the size of the set P we use in the algorithm. Similarly, we know that  $|P| << |\mathcal{O}_q|^2$ . Let  $\beta$  be the cost of Algorithm 2. It is easy to verify that the time complexity of MaxSum-Exact is  $O(n_1 \cdot |P| \cdot \beta)$ .

Next, we analyze  $\beta$ . The cost of lines 1-4 (Algorithm 2) is dominated by those of other parts in the algorithm. The cost of lines 5-6 is simply  $O(\log |\mathcal{O}| + |\mathcal{O}_q|)$  since we can issue three range queries and then perform an intersection on the query results. The cost of line 7 is  $O(|\psi| \cdot |\mathcal{O}_q|)$ . The cost of lines 8-11 is  $O(|\mathcal{O}'|^{|\psi|} \cdot |\psi|^2)$  since it enumerates at most  $O(|\mathcal{O}'|^{|\psi|})$  subsets S'' that cover  $\psi$  and each subset incurs a checking operation (line 10) whose cost is  $O(|\psi|^2)$  (since  $|S''| = O(|\psi|)$  and we can try all pairwise distances within S'' to do the checking). Thus,  $\beta$  is  $O(\log |\mathcal{O}| + |\mathcal{O}_q| + |\psi| \cdot |\mathcal{O}_q| + |\mathcal{O}'|^{|\psi|} \cdot |\psi|^2)$ . Note that  $|\mathcal{O}'| << |\mathcal{O}_q|$  (since  $\mathcal{O}'$  corresponds to a set of relevant objects in a small region),  $|\mathcal{O}_q| < |\mathcal{O}|$  and  $|\psi| \le |q.\psi| - 1$ .

In conclusion, the time complexity of MaxSum-Exact is  $O(n_1 \cdot |P| \cdot (\log |\mathcal{O}| +$ 

$$|\mathcal{O}_q| + |\psi| \cdot |\mathcal{O}_q| + |\mathcal{O}'|^{|\psi|} \cdot |\psi|^2).$$

## 2.3.2 Finding Approximate Solution

In this section, we propose a 1.375-factor approximate algorithm called *MaxSum-Appro* which is better than the best-known 2-factor approximate algorithm, Cao-Appro2.

Before we present MaxSum-Appro, we introduce the concept of "o-neighborhood feasible set". Given a query q and an object  $o \in \mathcal{O}$ , the o-neighborhood feasible set is defined to be the set containing o and all other objects each of which is the t-keyword nearest neighbor of o in D(q,d(o,q)) for each  $t \in q.\psi - o.\psi$ . For example, consider Figure 2.1. Suppose that the query  $q.\psi$  is  $\{t_1,t_2,t_3\}$ . Then, the  $o_1$ -neighborhood feasible set is  $\{o_1,o_2,o_3\}$  since  $q.\psi-o_1.\psi=\{t_1,t_3\},o_1$ 's  $t_1$ -keyword nearest neighbor in  $D(q,d(o_1,q))$  is  $o_2$  and  $o_1$ 's  $t_3$ -keyword nearest neighbor in  $D(q,d(o_1,q))$  is  $o_3$ . It could be easily verified that the o-neighborhood feasible set exists iff o is outside  $D(q,r_{min}-\delta)$  since an o-neighborhood feasible set is a feasible set.

In MaxSum-Appro, we only consider the o-neighborhood feasible sets for those objects o that are inside R(S) where S is a feasible set, and thus they always exist.

We present MaxSum-Appro in Algorithm 3. MaxSum-Appro is exactly Algorithm 1 by replacing Step 2 and Step 3 which are relatively expensive with the new efficient operation of finding the o-neighborhood feasible set which could be finished by issuing  $|q.\psi - o.\psi|$  NN queries.

**Theoretical Analysis.** Although the set S returned by the MaxSum-Appro algorithm might have a larger cost than the optimal set  $S_o$ , the difference is bounded.

**Theorem 2.3.2** MaxSum-Appro gives a 1.375-factor approximation for the MaxSum-CoSKQ problem.

#### Algorithm 3 Algorithm MaxSum-Appro

```
Require: query q and a set \mathcal{O} of objects
```

- 1:  $S \leftarrow N(q)$
- 2: while there is an "un-processed" relevant object o in R(S) do
- 3: // Step 1 (Query Distance Owner Finding)
- 4:  $o \leftarrow$  the nearest "un-processed" relevant object in R(S)
- 5: // Step 2 (o-Neighborhood Feasible Set Finding)
- 6:  $S' \leftarrow$  the *o*-neighborhood feasible set
- 7: **if** cost(S') < cost(S) **then**
- 8:  $S \leftarrow S'$
- 9: // Step 3 (Iterative Process)
- 10: mark o as "processed"
- 11: **return** *S*

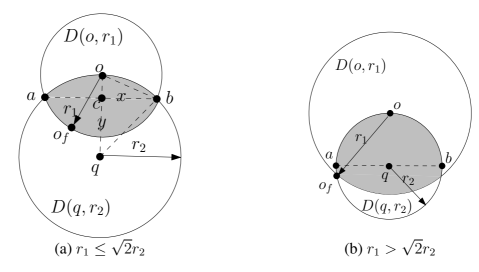


Figure 2.2: Illustration of the proof of Theorem 2.3.2

**Proof.** Let  $S_o$  be the optimal solution and S be the solution returned by MaxSum-Appro. Let o be the query distance owner of  $S_o$ . By Lemma 2.3.3, we know that o is in R(S). Besides, we can safely assume that o is a relevant object. Thus, there exists an iteration in MaxSum-Appro such that we process o (line 3) and thus we find its o-neighborhood feasible set denoted by S'.

Since S is the final solution returned by MaxSum-Appro, we know that  $cost(S) \le cost(S')$ . The remaining part of the proof shows that  $cost(S') \le 1.375 \cdot cost(S_o)$ .

Let  $o_f$  be the object in S' that is the farthest from o and  $r_1 = d(o_f, o)$ . Then, all objects in S' fall in  $D(o, r_1)$ . Let  $r_2 = d(o, q)$ . Since o is the query distance owner of S', we know that all objects in S' fall in  $D(q, r_2)$ . In summary, all objects in S' fall in

 $D(o, r_1) \cap D(q, r_2)$ .

Consider  $cost(S_o)$ . It could be verified by using a similar method for proving Lemma 2.3.1 that  $\max_{o_1,o_2\in S_o} d(o_1,o_2) \geq d(o,o_f)$ . Thus, we have  $cost(S_o) \geq r_2 + r_1$ .

In the following, we consider two cases on  $r_1$  according to whether there exists a line segment linking two points at the boundary of  $D(q, r_2)$  such that it has its length equal to  $2r_2$  (i.e., the diameter of  $D(q, r_2)$ ) and falls in  $D(o, r_1) \cap D(q, r_2)$ . Note that the boundary case happens when  $r_1 = \sqrt{2}r_2$  and there exists *exactly* one such segment.

Case 1:  $r_1 \leq \sqrt{2}r_2$ . We denote the intersection points between the boundaries of  $D(o,r_1)$  and  $D(q,r_2)$  by a and b, as shown in Figure 2.2(a). Let c be the intersection point between segment  $\overline{qo}$  and segment  $\overline{ab}$ . Let x=d(a,c)=d(b,c) and y=d(c,q). Since  $\triangle_{ocb}$  and  $\triangle_{qcb}$  are right-angled triangles, we know  $x^2+(r_2-y)^2=r_1^2$  and  $y^2+x^2=r_2^2$  by the *hypothesis theorem*. By solving these two equations, we obtain  $x=\sqrt{r_1^2-r_1^4/4r_2^2}$  and thus  $d(a,b)=2x=2\sqrt{r_1^2-r_1^4/4r_2^2}$ . In this case, it can be verified that  $\max_{o_1,o_2\in S'}d(o_1,o_2)\leq d(a,b)$  (since all objects in S' are in  $D(o,r_1)\cap D(q,r_2)$ , as shown in the shaded area of Figure 2.2(a)) and hence  $cost(S')\leq r_2+2\sqrt{r_1^2-r_1^4/4r_2^2}$ . Therefore,

$$\frac{cost(S')}{cost(S_o)} \le \frac{r_2 + 2\sqrt{r_1^2 - r_1^4/4r_2^2}}{r_2 + r_1} = 1 + \frac{2\sqrt{1 - r_1^2/4r_2^2} - 1}{r_2/r_1 + 1}$$

Let  $z=r_1/r_2$ . Thus,  $\frac{cost(S')}{cost(S_o)} \leq 1+\frac{2\sqrt{1-z^2/4}-1}{1/z+1}$ . Since  $r_1 \leq \sqrt{2}r_2$ , we have  $z \in (0,\sqrt{2}]^{-1}$ . We define  $f(z)=1+\frac{2\sqrt{1-z^2/4}-1}{1/z+1}$  on  $\{z|z\in(0,\sqrt{2}]\}$ . It could be verified that f(z) is monotonically increasing on (0,0.875) and is monotonically decreasing on  $(0.875,\sqrt{2}]$ . Thus,  $f(z)\leq f(0.875)<1.375$ . Therefore,

$$\frac{cost(S')}{cost(S_o)} \le f(z) \le 1.375$$

<sup>&</sup>lt;sup>1</sup>The interval  $(0, \sqrt{2}]$  does not include the boundary case where z=0 (i.e.,  $r_1=0$ ). In this case, we have  $cost(S')/cost(S_o)=1$ .

Case 2:  $r_1 > \sqrt{2}r_2$ . Let  $\overline{ab}$  be any segment linking two points at the boundary of  $D(q,r_2)$  which has its length equal to  $2r_2$  and falls in  $D(o,r_1)\cap D(q,r_2)$ . For illustration, consider Figure 2.2(b). That is,  $d(a,b)=2r_2$ . Similar to Case 1, it could be verified that  $\max_{o_1,o_2\in S'}d(o_1,o_2)\leq d(a,b)=2r_2$ . Thus,  $cost(S')\leq r_2+2r_2$ . Therefore,

$$\frac{cost(S')}{cost(S_o)} \le \frac{r_2 + 2r_2}{r_2 + r_1} = \frac{1+2}{1 + r_1/r_2} \le \frac{1+2}{1+\sqrt{2}} < 1.25$$

Thus, by combining Case 1 and Case 2, we have  $cost(S') \leq 1.375 \cdot cost(S_o)$ , which completes the proof.  $\blacksquare$ 

**Implementation and Time Complexity.** We also adopt the IR-tree built on  $\mathcal{O}$  to support the NN query and the range query.

Let  $n_1$  be the number of iterations in MaxSum-Appro (lines 2-10 in Algorithm 3) and  $\gamma$  be the cost of executing an iteration. Then, the time complexity of MaxSum-Appro is  $O(n_1 \cdot \gamma)$ . Note that  $\gamma$  is dominated by the step of finding the o-neighborhood feasible set (line 6) whose cost is bounded by  $O(|q.\psi| \cdot \log |\mathcal{O}|)$  (it issues at most  $|q.\psi - o.\psi|$  NN queries each of which takes  $O(\log |\mathcal{O}|)$  time). Thus,  $\gamma = O(|q.\psi| \cdot \log |\mathcal{O}|)$  where  $n_1 << |\mathcal{O}_q|$ . Note that the worst-case time complexity of MaxSum-Appro is  $O(|\mathcal{O}_q| \cdot |q.\psi| \cdot \log |\mathcal{O}|)$ , which is the same as that of Cao-Appro2.

# 2.4 Algorithms for Dia-CoSKQ

In this section, we propose two algorithms, Dia-Exact and Dia-Appro, for Dia-CoSKQ. Similarly, in this section, for clarity, we simply write  $cost_{Dia}(\cdot)$  as  $cost(\cdot)$  if the context of the cost function is clear.

### 2.4.1 Finding Optimal Solution

Interestingly, we can adopt the same MaxSum-Exact algorithm (Algorithm 1) by replacing the cost measurement from the maximum sum cost to the diameter cost. We call this algorithm Dia-Exact. The reason is that we can still use the query distance owner and the pairwise distance owners of a set S' to be found to find the optimal solution for Dia-CoSKQ. Next, we explain the reason in detail.

Consider the diameter cost. Given a set S' of objects in  $\mathcal{O}$ , we have  $cost(S') = \max_{o',o'' \in S' \cup \{o_q\}} d(o',o'')$ . Clearly, the (diameter) cost of a set S' can be dominated (or determined) by two pairwise distance owners of  $S' \cup \{o_q\}$  (not S' used in the maximum sum cost), which form a distance owner group (for Dia-CoSKQ). It is similar to the maximum sum cost of a set S' which is dominated by the query distance owner of S' and two pairwise distance owners of S'. But, there are two differences. The first difference is that the diameter cost is dominated by the pairwise distance owners only (without the query distance owner). The second difference is that the pairwise distance owners used for the diameter cost are based on the set  $S' \cup \{o_q\}$  instead of S'.

Based on the above observations, we *directly* adapt the distance owner-driven approach as follows. This approach maintains a variable S storing the best feasible set found so far. Initially, S is set to a feasible set. This involves three major steps.

- Step 1 (Pairwise Distance Owner Finding): Select two objects, o' and o'', in  $\mathcal{O}_q \cup \{o_q\}$  to take the roles of the pairwise distance owners of the set  $S' \cup \{o_q\}$  where S' is to be found. Note that o' and o'' form a distance owner group.
- Step 2 (Sub-Optimal Feasible Set Finding): Find a set S' of objects in  $\mathcal{O}_q$  such that the pairwise distance owners of  $S' \cup \{o_q\}$  are o' and o'' (if any), and update S with S' if cost(S') < cost(S).
- Step 3 (Iterative Step): Repeat Step 1 which finds another distance group, and continue with Step 2 until all distance owner groups have been traversed.

Interestingly, Step 1 which *originally* finds two objects to take the roles of the two pairwise distance owners *based on*  $S' \cup \{o_q\}$  can be *refined* to a number of sub-steps of finding two objects to take the roles of the two pairwise distance owners *based on* S' simply (not  $S' \cup \{o_q\}$ ) and finding an object to take the role of the query distance owner *based on* S'. This refinement can be explained by the following observation.

**Observation 1** Let S' be the feasible set. The pairwise distance owners of  $S' \cup \{o_q\}$  are either (1)  $o_q$  and the query distance owner of S' or (2) the pairwise distance owners of S'.

Suppose that o takes the role of the query distance owner of S' to be found, and  $o_1$  and  $o_2$  take the roles of the two pairwise distance owners of S'.

Observation 1 involves two cases. In Case (1) of Observation 1, we know that the pairwise distance owners of  $S' \cup \{o_q\}$  are  $o_q$  and the query distance owner o of S'. In this case, we deduce that  $d(o_1, o_2) \leq d(o, q) (= d(o, o_q))$ .

In Case (2) of Observation 1, we know that the pairwise distance owners of  $S' \cup \{o_q\}$  are the pairwise distance owners of S', say  $o_1$  and  $o_2$ . In this case,  $d(o,q) \le d(o_1,o_2)$ .

In conclusion, if we know that  $d(o_1, o_2) \leq d(o, q)$ , then  $o_q$  and o are the pairwise distance owners of  $S' \cup \{o_q\}$ . Otherwise,  $o_1$  and  $o_2$  are the pairwise distance owners of  $S' \cup \{o_q\}$ .

Thus, Step 1 can be refined with the following three sub-steps.

- Step I(a) (Query Distance Owner Finding): Select an object o in  $\mathcal{O}_q$  to take the role of the query distance owner of a set S' to be found.
- Step 1(b) (Pairwise Distance Owner Finding): Select two objects  $o_1$  and  $o_2$  in D(q, d(o, q)) to take the roles of the pairwise distance owners of the set S' to be found.

• Step I(c) (Pairwise Distance Owner Determination): If  $d(o,q) \ge d(o_1,o_2)$ , assign to o and  $o_q$  the roles of pairwise distance owners of  $S' \cup \{o_q\}$ ; otherwise, assign the roles to  $o_1$  and  $o_2$ .

With this refinement, the distance owner-driven approach still has its similar pruning features under the diameter cost. Specifically, Property 1 and Property 2 used for MaxSum-CoSKQ have their counterparts used for Dia-CoSKQ as Property 3 and Property 4, respectively.

**Property 3 (Pruning)** Let S' be a feasible set. If o is the query distance owner of S', then the two pairwise distance owners of  $S' \cup \{o_q\}$  are inside D(q, d(o, q)).

**Property 4 (Pruning)** Let S' be a feasible set, o be the query distance owner of S', and  $o_1$  and  $o_2$  be the two pairwise distance owners of  $S' \cup \{o_q\}$ . Then all objects in S' fall in  $D(q, d(o, q)) \cap D(o_1, d(o_1, o_2)) \cap D(o_2, d(o_1, o_2))$ .

Similar to the maximum sum cost, when the diameter cost is used, the object to be found in Step 1(a) is fetched based on the proximity to the query point q. The proximity is also related to the closest possible query distance owner (Lemma 2.3.1) and the farthest possible query distance owner (Lemma 2.3.2). It is easy to verify that Lemma 2.3.1 and Lemma 2.3.2 still hold when the cost measurement is changed from the maximum sum cost to the diameter cost. Thus, Lemma 2.3.3, which states that the *ring* is the region containing the query distance owners to be considered, still holds.

In summary, we present the algorithm for finding the optimal solution of Dia-CoSKQ in Algorithm 4 (which is quite similar to Algorithm 1) except that we need to determine the pairwise distance owner of  $S' \cup \{o_q\}$  (in Step 1(c)) which cannot be found in MaxSum-CoSKQ.

#### Algorithm 4 Algorithm Dia-Exact

```
Require: query q and a set \mathcal{O} of objects
 1: S \leftarrow N(q)
 2: while there is an "un-processed" relevant object o in R(S) do
        // Step 1(a) (Query Distance Owner Finding)
        o \leftarrow the nearest "un-processed" relevant object in R(S)
 4:
 5:
        // Step 1(b) (Pairwise Distance Owner Finding)
        D \leftarrow the q-disk with its radius equal to d(o, q)
 6:
        P \leftarrow \text{a set of all pairs } (o_1, o_2) \text{ where } o_1 \text{ and } o_2 \text{ are in } D
 7:
        for each (o_1, o_2) \in P in ascending order of d(o_1, o_2) do
 8:
           // Step 1(c) (Pairwise Distance Owner Determination)
 9:
           if d(o,q) \ge d(o_1,o_2) then o' \leftarrow o; o'' \leftarrow o_a
10:
           else o' \leftarrow o_1; o'' \leftarrow o_2
11:
           // Step 2 (Sub-Optimal Feasible Set Finding)
12:
           if there exists a feasible set S' in D which is (o, o', o'')-owner consistent then
13:
              \begin{array}{c} \textbf{if } cost(S') < cost(S) \textbf{ then} \\ S \leftarrow S'; \textbf{ break} \end{array}
14:
15:
        // Step 3 (Iterative Process)
16:
        mark o as "processed"
17:
18: return S
```

**Theorem 2.4.1** Dia-Exact returns a feasible set with the smallest cost for the Dia-CoSKQ problem.

**Proof.** Let  $S_o$  be one of the feasible set with the smallest cost. Let o be the query distance owner of  $S_o$ , and let  $o_1$  and  $o_2$  be the two pairwise distance owners of  $S_o$ . First, o is inside R(S) (Lemma 2.3.3). Thus, there exists an iteration where o is processed. When o is processed, pair  $(o_1, o_2)$  must be included in P (Property 3). There are two cases. Case 1:  $d(o,q) \geq d(o_1,o_2)$ . In this case, any feasible set S' that is  $(o,o,o_q)$ -owner consistent is retrieved and used to update S (there must exist some since  $S_o$  is  $(o,o,o_q)$ -owner consistent). Thus, the resulting S has its cost equal to  $d(o,q) = cost(S_o)$ . Case 2:  $d(o,q) < d(o_1,o_2)$ . In this case, any feasible set S' that is  $(o,o_1,o_2)$ -owner consistent is retrieved and used to update S (there must exist some since  $S_o$  is  $(o,o_1,o_2)$ -owner consistent). Thus, the resulting S has its cost equal to  $d(o_1,o_2) = cost(S_o)$ . In either case, S will not be updated anymore since it has the smallest cost (i.e.,  $cost(S_o)$ ) and thus it is the final output.

Same as Section 2.3.1, in Dia-Exact, we have the self-iteration computation strat-

egy and the cross-iteration computation strategy.

**Self-Iteration Computation Strategy:** Consider an iteration where the query distance owner for this iteration is o. We can use the same mechanism described in Section 2.3.1 after  $d_{min}$  and  $d_{max}$  are updated from  $d(o,q) - \min\{d(o_1,q), d(o_2,q)\}$  and cost(S) - d(o,q) to d(o,q) and cost(S), respectively. All pruning properties still hold.

Note that  $d_{min}$  (which is originally set to  $d(o,q) - \min\{d(o_1,q),d(o_2,q)\}$  in MaxSum-CoSKQ) is based on the triangle inequality (Lemma 2.3.4), which means that it can be used for pruning in both MaxSum-CoSKQ and Dia-CoSKQ. However, in Dia-CoSKQ,  $d_{min}$  can be updated to a tighter value as d(o,q) since all pairs with their pairwise distances smaller than d(o,q) cannot take the roles of the pairwise distance owners of  $S' \cup \{o_q\}$ .

Cross-Iteration Computation Strategy: We use the same information reuse techniques as in Section 2.3.1 for Dia-Exact since the updated  $d_{min}$  (i.e., d(o,q)) is monotonically increasing and the updated  $d_{max}$  (i.e., cost(S)) is monotonically decreasing with more iterations. Thus, the pairs pruned in P at the previous iterations need not be considered in the later iterations.

**Time Complexity.** It could be verified that the time complexity of Dia-Exact is the same as that of MaxSum-Exact.

## 2.4.2 Finding Approximate Solution

In this section, we propose a  $\sqrt{3}$ -factor approximate algorithm which is exactly the same as Algorithm 3 but the cost measurement used is the diameter cost. This algorithm is called *Dia-Appro*.

**Theoretical Analysis.** Although the set S returned by Dia-Appro may have a larger cost than the optimal set  $S_o$ , it has an approximate factor of  $\sqrt{3}$ .

**Theorem 2.4.2** Dia-Appro gives a  $\sqrt{3}$ -factor approximation for the Dia-CoSKQ problem.

**Proof.** We use the same notations as defined in the proof of Theorem 2.3.2.

Consider  $cost(S_o)$ . Similar to the proof of Theorem 2.3.2, we have  $\max_{o'_1,o'_2\in S_o}d(o'_1,o'_2)\geq d(o,o_f)=r_1$ . Recall that  $\max_{o'\in S_o}d(o',q)=d(o,q)=r_2$ . As a result, we have  $cost(S_o)=\max\{\max_{o'\in S_o}d(o',q),\max_{o'_1,o'_2\in S_o}d(o'_1,o'_2)\}\geq \max\{r_2,r_1\}$ .

According to the Dia-Appro algorithm, we have  $cost(S) \leq cost(S')$ . The remaining part of the proof shows that  $cost(S') \leq \sqrt{3} \cdot cost(S_o)$  which further implies  $cost(S) \leq \sqrt{3} \cdot cost(S_o)$ .

Same as the proof of Theorem 2.3.2, we consider two cases of  $r_1$ .

Case 1:  $r_1 \leq \sqrt{2}r_2$ . This case corresponds to Figure 2.2(a). It can be verified that  $\max_{o'_1,o'_2\in S'}d(o'_1,o'_2)\leq d(a,b)=2\sqrt{r_1^2-r_1^4/4r_2^2}$  (since all objects in S' fall in  $D(o,r_1)\cap D(q,r_2)$  as shown by the shaded area). Recall  $\max_{o'\in S'}d(o,q)=r_2$ . As a result, we have  $cost(S')\leq \max\{r_2,2\sqrt{r_1^2-r_1^4/4r_2^2}\}$ .

We further consider three sub-cases under Case 1 based on the relationship among  $r_1$ ,  $r_2$  and  $2\sqrt{r_1^2 - r_1^4/4r_2^2}$ .

Case 1(a):  $r_1 \leq \sqrt{2-\sqrt{3}}r_2$ . In this case, we have  $r_2 > r_1$  and  $r_2 \geq 2\sqrt{r_1^2-r_1^4/4r_2^2}$ . Thus,  $cost(S_o) \geq \max\{r_2,r_1\} = r_2$  and  $cost(S') \leq \max\{r_2,2\sqrt{r_1^2-r_1^4/4r_2^2}\} = r_2$  Therefore,

$$\frac{cost(S')}{cost(S_o)} \le \frac{r_2}{r_2} = 1$$

Case 1(b):  $\sqrt{2-\sqrt{3}}r_2 < r_1 \le r_2$ . In this case, we have  $r_2 \ge r_1$  and

 $2\sqrt{r_1^2 - r_1^4/4r_2^2} > r_2$ . Thus,  $cost(S_o) \ge r_2$  and  $cost(S') \le 2\sqrt{r_1^2 - r_1^4/4r_2^2}$ . Therefore,

$$\frac{cost(S')}{cost(S_o)} \le \frac{2\sqrt{r_1^2 - r_1^4/4r_2^2}}{r_2} = \sqrt{4(\frac{r_1}{r_2})^2 - (\frac{r_1}{r_2})^4}$$
 (2.4)

Note that function  $f(z)=\sqrt{4z^2-z^4}$  is monotonically increasing on  $(\sqrt{2-\sqrt{3}},1]$ . Since  $\frac{r_1}{r_2}\in(\sqrt{2-\sqrt{3}},1]$ , Thus, we have

$$\frac{cost(S')}{cost(S_o)} \le \sqrt{4(1)^2 - (1)^4} = \sqrt{3}$$

Case 1(c):  $r_2 < r_1 \le \sqrt{2}r_2$ . In this case, we have  $r_2 < r_1$  and  $2\sqrt{r_1^2 - r_1^4/4r_2^2} > r_2$ . Thus,  $cost(S_o) \ge r_1$  and  $cost(S') \le 2\sqrt{r_1^2 - r_1^4/4r_2^2}$ . Therefore,

$$\frac{cost(S')}{cost(S_o)} \le \frac{2\sqrt{r_1^2 - r_1^4/4r_2^2}}{r_1} = \sqrt{4 - (r_1/r_2)^2} < \sqrt{3}$$

Case 2:  $r_1 > \sqrt{2}r_2$ . This case corresponds to Figure 2.2 (b). In this case,  $d(a,b) = 2r_2$ . Similar to Case 1, we have  $\max_{o_1,o_2 \in S'} d(o_1,o_2) \le d(a,b) = 2r_2$ . Therefore,

$$\frac{cost(S')}{cost(S_o)} \le \frac{\max\{r_2, 2r_2\}}{\max\{r_1, r_2\}} = \frac{2r_2}{r_1} < \frac{2r_2}{\sqrt{2}r_2} = \sqrt{2}$$

In view of above discussion, we know that  $cost(S')/cost(S_o) \leq \sqrt{3}$ , which completes the proof.  $\blacksquare$ 

**Time Complexity.** Since Dia-Appro is identical to MaxSum-Appro except that Dia-Appro adopts a different cost measurement, Dia-Appro has the same complexity as MaxSum-Appro.

### 2.4.3 Adaptions of Existing Solutions

In this section, we adapt the existing solutions in [11], which are originally designed for MaxSum-CoSKQ, for Dia-CoSKQ.

Cao-Exact. Cao-Exact is a best-first search method based on the object space and thus its applicability is independent of the cost measurement used in the CoSKQ problem. Therefore, Cao-Exact can be directly applied to Dia-CoSKQ by replacing the cost measurement with the diameter cost. However, due to its prohibitively huge search space, Cao-Exact is not scalable to large datasets.

**Cao-Appro1 & Cao-Appro2.** We can directly adopt Cao-Appro1 and Cao-Appro2 for Dia-CoSKQ by replacing the maximum sum cost with the diameter cost.

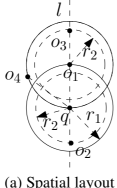
According to [11], the approximation factors of Cao-Appro1 and Cao-Appro2 are 3 and 2, respectively, for MaxSum-CoSKQ. In the following, we prove that both Cao-Appro1 and Cao-Appro2 give 2-factor approximations for Dia-CoSKQ.

**Lemma 2.4.1** Cao-Approl and Cao-Appro2 give 2-factor approximations for Dia-CoSKQ.

**Proof.** First, we prove that the approximation ratio of Cao-Appro1 is 2.

Let S be the set returned by Cao-Appro1 and  $S_o$  be the optimal set. Let  $o_f$  be the object in S that is the farthest from q, i.e.,  $d(o_f,q) = \max_{o \in S} d(o,q)$ . First, we have  $cost(S_o) \geq d(o_f,q)$ . Second, for any two objects  $o_1$  and  $o_2$  in S, we have  $d(o_1,o_2) \leq d(o_1,q) + d(o_2,q) \leq 2 \cdot d(o_f,q)$  by the *triangle inequality*. Therefore,  $cost(S) \leq \max\{d(o_f,q), 2 \cdot d(o_f,q)\} = 2 \cdot d(o_f,q)$ . As a result, we know  $cost(S)/cost(S_o) \leq 2$ .

Since the solution returned by Cao-Appro2 is no worse than that returned by Cao-Appro1, the approximation ratio of Cao-Appro2 is also bounded by 2.



object	keywords		
q	$t_1, t_2$		
$o_1$	$t_1$		
$o_2$	$t_2$		
$o_3$	$t_2$		
$o_4$	$t_2$		

(a) Spatial layout

(b) Keyword information

Figure 2.3: A problem instance for Cao-Appro2

Furthermore, we show that Cao-Appro2 cannot provide better error guarantees by constructing a problem instance where the approximation ratio of Cao-Appro2 is infinitely close to 2.

The problem instance is shown in Figure 2.3. In Figure 2.3(a), we have four objects,  $o_1$ ,  $o_2$ ,  $o_3$  and  $o_4$ .  $o_1$ ,  $o_2$  and  $o_3$  are located at the boundary of  $D(q, r_1)$ ,  $D(q, r_2)$  and  $D(o_1, r_2)$ , respectively.  $r_2 = r_1 - \delta$  ( $\delta > 0$ ). Besides,  $q, o_1, o_2$  and  $o_3$  are on the same vertical line l.  $o_4$  is on the boundary of  $D(o_1, r_1)$  and outside  $D(q, r_1)$ . In addition,  $d(o_4,q)=r_1+\delta$ . The keyword information of these objects and the query are shown in Figure 2.3(b).

Given the above problem instance, Cao-Appro2 works as follows. First, it invokes the Cao-Appro1 algorithm which returns  $S_1 = \{o_1, o_2\}$  as the first candidate of the approximate solution whose cost is equal to  $d(o_1, o_2) = r_1 + r_2$ . Thus,  $o_1$  is the farthest object in  $S_1$  from q and  $t_1$  is the keyword that is covered by  $o_1$  but not by any other objects in  $S_1$ . As a result, Cao-Appro2 would invoke Cao-Appro1 for each object containing  $t_1$ . In this problem instance, since only  $o_1$  contains  $t_1$ , Cao-Appro2 invokes Cao-Appro1 at  $o_1$  only and  $S_2 = \{o_1, o_3\}$  would be returned as the second candidate of the approximate solution whose cost is equal to  $d(o_3, q) = r_1 + r_2$ . Since Cao-Appro2 obtains two candidates  $S_1$  and  $S_2$  with the same cost, it returns any of them, say  $S_2$ , as the final approximate solution S. However, the optimal solution  $S_o$  of this problem

instance is  $\{o_1, o_4\}$  with its cost equal to  $d(o_4, q) = r_1 + \delta$ . Therefore,

$$\frac{cost(S_2)}{cost(S_o)} = \frac{r_1 + r_2}{r_1 + \delta} = 2 - \frac{3}{r_1/\delta + 1}$$

When  $\delta$  approaches 0, the ratio approaches 2.

Thus, among all known approximate algorithms for Dia-CoSKQ, our Dia-Appro provides the best constant-factor approximation.

# 2.5 Empirical Studies

## 2.5.1 Experimental Set-up

**Datasets.** We used the real datasets adopted in [11], namely Hotel, Web and GN. Dataset Hotel corresponds to a set of hotels in the U.S. (www.allstays.com), each of which is associated with its location and a set of words that describe the hotel (e.g., restaurant and pool). Dataset Web was created from two real datasets. The first one, named WEBSPAMUK2007<sup>2</sup>, corresponds to a set of web documents. The second one is a set of spatial objects, named TigerCensusBlock<sup>3</sup>, which corresponds to a set of census blocks in Iowa, Kansas, Missouri and Nebraska. Specifically, Web consists of the spatial objects in TigerCensusBlock, each of which is associated with a document randomly selected from WEBSPAMUK2007. Dataset GN was collected from the U.S. Board on Geographic Names (geonames.usgs.gov). Each object in GN is a 2D location which is associated with a set of keywords describing it (e.g., a geographic name like valley).

**Query Generation.** Given a dataset  $\mathcal{O}$  and a positive integer k, we generated a query q with the size of its keyword set equal to k as in [11]. For the  $q.\lambda$  part, we randomly

<sup>&</sup>lt;sup>2</sup>http://barcelona.research.yahoo.net/webspam/datasets/uk2007

<sup>&</sup>lt;sup>3</sup>http://www.rtreeportal.org

Statistics	GN	Web	Hotel
Number of objects	1,868,821	579,727	20,790
Number of unique words	222,409	2,899,175	602
Number of words	18,374,228	249,132,883	80,845

Table 2.1: Real datasets (for CoSKQ)

picked a location from the data space of  $\mathcal{O}$ . For the  $q.\psi$  part, we first sorted all the keywords that are associated with the objects in  $\mathcal{O}$  in descending order of their frequencies and then randomly picked k keywords among all keywords each of which has its *percentile rank* within range [10,40] by default. Note that in this way, each of the keywords in  $q.\psi$  has a relatively high frequency.

**Algorithms.** For MaxSum-CoSKQ, we consider 2 exact algorithms, namely MaxSum-Exact and Cao-Exact, and 3 approximate algorithms, namely MaxSum-Appro, Cao-Appro1 and Cao-Appro2. For Dia-CoSKQ, we consider 2 exact algorithms, namely Dia-Exact and Cao-Exact (the adaption), and 3 approximate algorithms, namely Dia-Appro, Cao-Appro1 and Cao-Appro2. All algorithms were implemented in C/C++.

Our experiments were conducted on a Linux platform with a 2.66GHz machine and 4GB RAM.

## 2.5.2 Experimental Results

We consider 2 measurements, the running time and the approximation ratio (for approximate algorithms only). For each set of settings, we generated 50 queries, ran the algorithms with each of these 50 queries, and averaged the experimental measurements.

#### **Experiments for MaxSum-CoSKQ**

**Effect of**  $|q.\psi|$ . We generated 5 types of queries with different values of  $|q.\psi|$ . The values we used are 3, 6, 9, 12 and 15. The results on the dataset GN are shown in

Figure 2.4. According to Figure 2.4(a), our MaxSum-Exact is faster than Cao-Exact by 1-3 orders of magnitude. When  $|q.\psi|$  increases, the running time gap between MaxSum-Exact and Cao-Exact increases. Besides, MaxSum-Appro and Cao-Appro2 have comparable running time, which verified our theoretical analysis that MaxSum-Appro and Cao-Appro2 have the same worst-case time complexity. Cao-Appro1 runs the fastest due to its simplicity. According to Figure 2.4(b), the approximation ratio of our MaxSum-Appro algorithm is near to 1, which shows that the accuracy of MaxSum-Appro is extremely high in practical. We note here that the approximation ratio in the figure corresponds to the average over 50 queries, among which, the approximation ratio of MaxSum-Appro is exactly 1 for most queries (e.g., more than 45). As a result, the approximation ratio of MaxSum-Appro in the figures is always near to 1. Consistent to our theoretical results, the approximation ratios of Cao-Appro1 and Cao-Appro2 are larger than that of MaxSum-Appro.

We have similar results on Web (Figure 2.5) and Hotel (Figure 2.6).

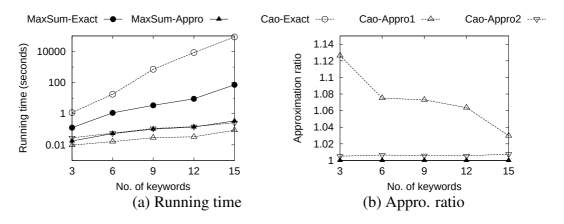


Figure 2.4: Effect of  $|q.\psi|$  (GN, MaxSum-CoSKQ)

Effect of average  $|o.\psi|$ . Our experiments were based on dataset Hotel whose average size of a keyword set of an object  $(|o.\psi|)$  is nearly 4 (i.e., 80,845/20,790). We generated a set of several datasets based on dataset Hotel such that the average sizes (i.e., average  $|o.\psi|$ 's) are equal to  $4 \cdot i$  for some integers i. To generate a dataset with its average  $|o.\psi|$  equal to  $4 \cdot i$ , we proceed with i-1 rounds. At each round, for each object o in

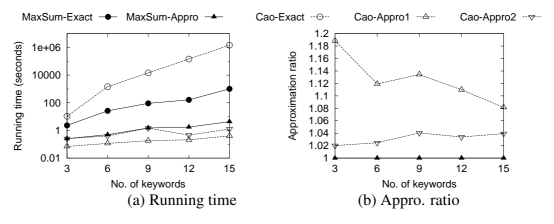


Figure 2.5: Effect of  $|q.\psi|$  (Web, MaxSum-CoSKQ)

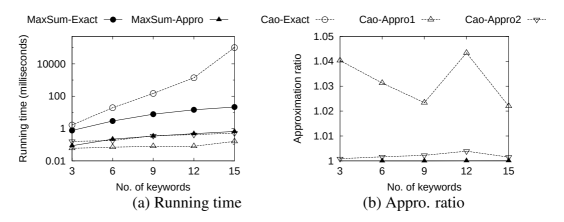


Figure 2.6: Effect of  $|q.\psi|$  (Hotel, MaxSum-CoSKQ)

dataset Hotel, we randomly pick another object o' and update  $o.\psi$  to be  $o.\psi \cup o'.\psi$ . It could be verified that the average  $|o.\psi|$  of the resulting dataset is nearly  $4 \cdot i$ . In our experiments, we vary i by choosing one of the values in  $\{1, 2, 4, 6, 8, 10\}$ . Note that i=1 means that the resulting dataset is exactly dataset Hotel.

The results are shown in Figure 2.7. According to Figure 2.7(a), the running times of all algorithms increase when the average  $|o.\psi|$  increases. The reason is that when the average  $|o.\psi|$  increases, the number of relevant objects  $(|\mathcal{O}_q|)$  in the dataset would probably increase, which further affects the running times of the algorithms. Since all algorithms except for Cao-Appro1 have their time complexities involving  $|\mathcal{O}_q|$ . Cao-Appro1, though has its time complexity independent of  $|\mathcal{O}_q|$ , has its NN queries affected by  $|\mathcal{O}_q|$ : the larger  $|\mathcal{O}_q|$  is, the more expensive the NN query would probably be. Besides, it is worth mentioning that when the average  $|o.\psi|$  increases, the increase

rate of the running time of Cao-Exact is significantly larger than those of the other algorithms including MaxSum-Exact. This is because Cao-Exact is based on the search space of the set of all possible feasible sets whose size increases rapidly with  $|\mathcal{O}_q|$  ( $|\mathcal{O}_q|^{|q,\psi|}$ ). Thus, Cao-Exact is not scalable on datasets with a large average  $|o.\psi|$ . According to Figure 2.7(b), the average  $|o.\psi|$  has no obvious trend on the approximation ratios of the approximate algorithms. Besides, MaxSum-Appro with its approximation ratio near to 1 always keeps its accuracy superiority over other approximate algorithms.

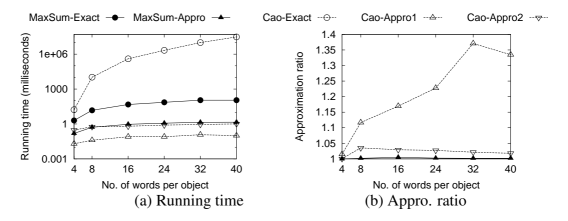


Figure 2.7: Effect of average  $|o.\psi|$  (MaxSum-CoSKQ)

Scalability Test. We conducted a scalability test on the algorithms with 5 synthetic datasets with their sizes varying from 2M to 10M. The synthetic datasets were generated from a smaller dataset GN. To generate a dataset  $\mathcal{O}$  with its size equal to n, we first inserted all the objects from dataset GN into  $\mathcal{O}$  and then repeatedly created objects in  $\mathcal{O}$  such that  $\mathcal{O}$  has a similar spatial distribution as dataset GN until  $|\mathcal{O}| = n$ . For each newly created object o in  $\mathcal{O}$ , we randomly pick a document from WEBSPAMUK2007 and use it as  $o.\psi$ .

The results are shown in Figure 2.8(a), where we do not show the running time of the algorithm if it runs more than 10 days or out of memory. According to these results, both our exact algorithm (MaxSum-Exact) and our approximate algorithm (MaxSum-Appro) are scalable to large datasets with millions of objects. For example, in a dataset with size equal to 10M, MaxSum-Exact ran less than 100s and MaxSum-Appro ran in

real-time. In contrast, Cao-Exact is not scalable. In particular, in our experiments, Cao-Exact took more than 1 day on a dataset with size equal to 6M and it took more than 10 days on a dataset with size equal to 8M.

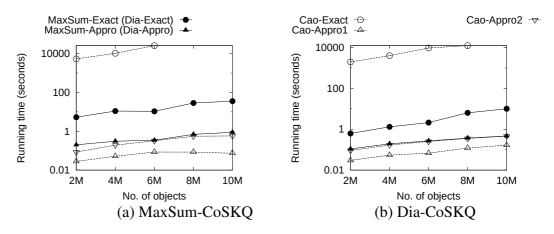


Figure 2.8: Scalability Test (CoSKQ)

#### **Experiments for Dia-CoSKQ**

**Effect of**  $|q.\psi|$ . The results on dataset GN is shown in Figure 2.9. According to Figure 2.9(a), our Dia-Exact is faster than Cao-Exact by 1-4 orders of magnitude. When  $|q.\psi|$  increases, the running time gap between Dia-Exact and Cao-Exact increases. Besides, Dia-Appro and Cao-Appro2 have comparable running times. According to Figure 2.9(b), similar to MaxSum-Appro (for MaxSum-CoSkQ), the approximation ratio of Dia-Appro is near to 1 (for Dia-CoSKQ), which is better than those of Cao-Appro1 and Cao-Appro2.

The results on datasets Web and Hotel are similar and thus they are omitted here due to the page limit.

**Effect of Average**  $|o.\psi|$ . Similar to the experiments for MaxSum-CoSKQ, we generated a set of datasets by varying their average  $|o.\psi|$  values. The results are shown in Figure 2.10. According to Figure 2.10(a), when the average  $|o.\psi|$  increases, the running time of Cao-Exact increases significantly while the running times of other al-

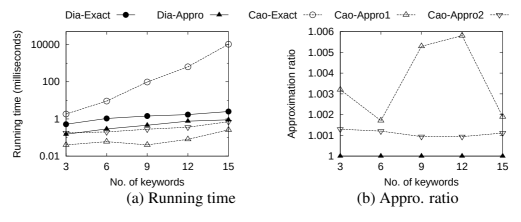


Figure 2.9: Effect of  $|q.\psi|$  (GN, Dia-CoSKQ)

gorithms are only slightly affected. This is similar to the case for MaxSum-CoSKQ and the explanation for MaxSum-CoSKQ as we discussed previously could be applied here for Dia-CoSKQ. According to Figure 2.10(b), the average  $|o.\psi|$  value has no obvious trend on the accuracy of the approximate algorithms.

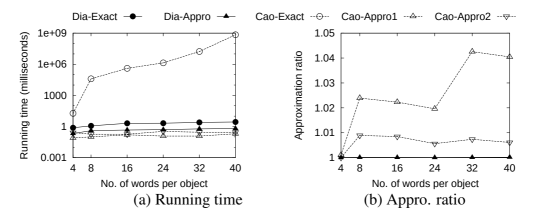


Figure 2.10: Effect of average  $|o.\psi|$  (Dia-CoSKQ)

**Scalability Test.** We conducted a scalability test on the algorithms for Dia-CosKQ with the same synthetic datasets used in the scalability test for MaxSum-CoSKQ.

The results are shown in Figure 2.8(b), where we do not show the running time of the algorithm if it runs more than 10 days or out of memory. According to these results, both our exact algorithm (Dia-Exact) and our approximate algorithm (Dia-Appro) are scalable to large datasets with millions of objects. In contrast, Cao-Exact is not scalable to large datasets.

**Conclusion:** MaxSum-Exact (Dia-Exact) runs faster than Cao-Exact by several orders of magnitude for MaxSum-CoSKQ (Dia-CoSKQ). Besides, MaxSum-Exact (Dia-Exact) is scalable in terms of  $|\mathcal{O}|$  as well as the average  $|o.\psi|$  but Cao-Exact is not. Our MaxSum-Appro (Dia-Appro) has a better accuracy while having comparable running time as those existing approximate algorithms.

## 2.6 Related Work

Many types of spatial keyword query have been proposed in the literature. Most of them are different from CoSKQ studied in this thesis since they use a single object to cover all keywords specified in the query but CoSKQ uses multiple objects collectively for the same purpose. We review these spatial keyword queries as follows.

A spatial keyword top-k query [19] finds top-k objects where the ranking function takes both the spatial proximity and the textual relevance of the objects into consideration. This branch includes [19, 56, 47] (Euclidean space), [57] (road networks), [58, 20] (trajectory databases), and [68] (moving objects). A common technique shared by these studies is to design a hybrid indexing structure, which captures both the spatial proximity and the textual information of the objects. The IR-tree adopted by us for NN queries and range queries was proposed in [19].

A spatial keyword k-NN query [27] finds the k-NNs from the query location, each of which contains the set of keywords specified in the query. That is, unlike the keywords in the spatial keyword top-k queries, which are used as a *soft* constraint, the keywords in the spatial keyword k-NN queries are used as a *hard* constraint. This branch includes [27, 12, 67].

A spatial keyword range query [63, 75, 18] takes a region and a set of keywords as input and finds the objects each of which falls in the region and contains the set of

keywords. Same as the spatial keyword k-NN queries, the keywords are used as a hard constraint. Usually, they combine a spatial index (e.g., R-tree and Space Filling Curve (SFC)) and a textual index (e.g., inverted file) for query processing.

A spatial keyword reverse top-k query [49] finds the set of objects whose spatial keyword top-k query results include the query. Note that in this case, an object which consists of a location and a set of keywords could be regarded as a query which also consists of a location and a set of keywords and vice versa.

An mCK query [71, 72] is a spatial keyword query that is very similar to CoSKQ. An mCK query takes m keywords as input and finds m objects with the smallest di-ameter that cover the m keywords specified in the query. Though both the mCK query and CoSKQ use a set of objects for covering a set of keywords collectively, they are different. In the context of an mCK query, it is assumed that each object is associated with a single keyword while in the context of CoSKQ, each object is associated with a set of multiple keywords. Besides, an mCK query only takes a set of keywords as input while our CoSKQ query takes not only a set of keywords but also a query location as an input.

CoSKQ was first studied in [11]. Under the maximum sum cost function, as we described, [11] proposed an exact algorithm and two approximate algorithms. However, the exact algorithm is not scalable to large datasets and the two approximate algorithms cannot guarantee near-to-optimal solutions. In this thesis, we propose an efficient exact algorithm and an approximate algorithm with better approximate factor for MaxSum-CoSKQ. Besides, in this thesis, we also propose another cost function called the diameter function which is new and has not been studied in [11].

## 2.7 Conclusion

In this chapter, we study two types of the CoSKQ problem, namely MaxSum-CoSKQ and Dia-CoSKQ. MaxSum-CoSKQ is a CoSKQ problem using the existing maximum sum cost, which is NP-hard. We design two algorithms for MaxSum-CoSKQ, MaxSum-Exact and MaxSum-Appro. MaxSum-Exact is an exact algorithm which significantly outperforms its existing competitor in terms of both efficiency and scalability and MaxSum-Appro is an approximate algorithm which improves the best-known constant approximation factor from 2 to 1.375. We also propose a new cost function and the CoSKQ problem using this function is Dia-CoSKQ. We design two algorithms for Dia-CoSKQ, Dia-Exact and Dia-Appro. Dia-Exact is an exact algorithm while Dia-Appro is a  $\sqrt{3}$ -factor approximate algorithm. Extensive experiments were conducted which verified our theoretical findings and algorithms.

## CHAPTER 3

# WORST-CASE OPTIMIZED SPATIAL MATCHING

## 3.1 Introduction

Bichromatic reverse nearest neighbor (BRNN) queries have been studied extensively [38, 41, 65]. Let P be a set of service-providers and O be a set of customers. A BRNN query is to find which customers in O are "interested" in a given service-provider in P. However, BRNN queries lack the consideration of the capacities of service-providers and the demands of customers. In order to address this issue, some spatial matching problems [66, 62, 61] have been proposed which assign service-providers to customers with the above consideration.

In some real-life applications like hospital allocation, a common goal is to minimize the *maximum* distance (or cost) between a hospital and a residential estate served by this hospital. For example, in the Hong Kong ambulance service, the minimized maximum cost is about 12 minutes (driving distance) [1].

To illustrate, we go through a toy example as shown in Figure 3.1. In Figure 3.1(a), P contains three hospitals  $p_1$ ,  $p_2$  and  $p_3$  and O contains three residential estates  $o_1$ ,  $o_2$  and  $o_3$ . Figure 3.1(b) shows all pairwise distances between P and O. For the sake of illustration, suppose that the *capacity* of each hospital p in P is 1, which means that the greatest amount of the service given by p is 1, and the *demand* of each residential estate o in O is also 1, which means that the amount of the service requested by o is 1. In this case, each hospital can serve at most one residential estate. In order to minimize the maximum distance between a hospital and the residential estate served

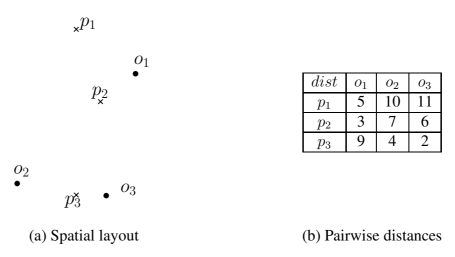


Figure 3.1: A running example (for SPM-MM)

by this hospital, we form an *assignment* between P and O as shown in Figure 3.2(a). In this assignment,  $p_1, p_2$  and  $p_3$  serve  $o_1, o_3$  and  $o_2$ , respectively. If p serves o, we draw a line between p and o in the figure. The number next to the line is called the *matching distance* between p and o which corresponds to the Euclidean distance between p and o. In this assignment, the *maximum matching distance* (mmd) is equal to 6. Besides, we cannot find any other assignment which satisfies the service demand of each customer and has its p and p smaller than 6. Thus, 6 is the p and p an

In this thesis, we propose a new problem called <u>SPatial Matching for Minimizing Maximum matching distance</u> (SPM-MM). Given a set P of service-providers each of which has a capacity and a set O of customers each of which has a demand, the SPM-MM problem is to assign the service-providers in P to the customers in O with the consideration of the capacities of the service-providers such that the demand of each customer in O is satisfied and the maximum matching distance (i.e. mmd) is minimized.

SPM-MM has extensive applications in matching between two sets of objects where the *worst-case* cost should be minimized. The notions of "service-provider" and "customer" in SPM-MM are general and can have alternative semantics in different (even non-geographic) applications. One such application is the allocation problem

between emergency facilities and users. Hospitals, fire stations and police stations are some examples of emergency facilities and residential estates and commercial areas are some examples of users. Logistics, data warehouse allocation and mail delivery are some applications with non-emergency facilities. *Profile matching* [66] is another application where we want to match "items" (regarded as service-providers) with "customers" such that the worst-case dissatisfactory rate among all customers is minimized.

It turns out that SPM-MM reduces to be a classical problem in computer science,  $Bottleneck\ Matching\ Problem\ (BMP)\ [32]$ , when each service-provider has its capacity equal to 1 and each customer has its demand equal to 1 as well. Given two sets of n objects, A and B, and the cost of matching each object in A with each object in B, the BMP problem is to find the  $perfect\ matching$  with the smallest cost among all perfect matchings between A and B where the cost of a perfect matching M is defined to be the greatest cost of matching an object from A and an object from B in A. It can be verified that SPM-MM becomes BMP when |P|=|O|, each service-provider  $p\in P$  (customer  $o\in O$ ) has its capacity (demand) equal to 1, and the distance between p and o is used as the cost of matching p with o for each  $p\in P$  and each  $o\in O$ . [8] provides a comprehensive study on existing solutions of BMP, among which, the Threshold algorithm is the fastest.

No existing algorithms can be used to solve the SPM-MM problem. Firstly, the algorithms for BMP cannot be used *directly* for SPM-MM since in SPM-MM, the capacities/demands could be arbitrary positive integers. Besides, we will show that an *adapted* version of the *Threshold* algorithm, which is originally designed for BMP, is not scalable for SPM-MM. Secondly, the solutions for all existing spatial matching problems cannot be used for SPM-MM. To illustrate this, we first give a brief background of these problems. Two major types of spatial matching problems have been studied. The first one [66] aims to find the *fair* assignment between *P* and *O* which

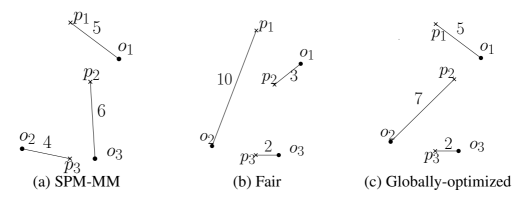


Figure 3.2: Spatial matching problems, SPM-MM

is to assign to each customer the nearest service-provider that has not been exhausted of serving other *closer* customers. Figure 3.2(b) shows the fair assignment between P and O whose mmd is equal to  $10 \ (> 6)$ . The second one is to find the globally optimized assignment between P and O which guarantees that each customer's service demand is satisfied and the overall matching cost is minimized. Figure 3.2(c) shows the globally optimized assignment between P and O whose mmd is equal to P (> 6).

In this thesis, we design two algorithms for the SPM-MM problem. The first one is called *Threshold-Adapt* and the second one is called *Swap-Chain*. *Threshold-Adapt* is an algorithm which shares a similar idea as *Threshold* which is originally designed for BMP. Unfortunately, *Threshold-Adapt* is not scalable to large datasets due to its high time/space complexity. *Swap-Chain* is an algorithm which is scalable and runs faster than *Threshold-Adapt* by orders of magnitude by using the concept of finding a series of elements where every two adjacent elements are "close" to each other for re-matching. The operation of finding a "close" element from another element can be implemented efficiently by spatial queries.

It is worth mentioning that our proposed algorithms are not limited to the Euclidean space. In fact, they can also be adapted to non-metric space (with a certain sacrifice of efficiency). For example, our algorithms can be adapted to settle the SPM-MM problem in Figure 3.1, even if the distance between a hospital and a residential estate is their road-network distance. The discussion on how to adapt our techniques to non-

metric space is given in Section 3.6.

We summarize our main contributions as follows. Firstly, to the best of our knowledge, we are the first to propose the SPM-MM problem, which has extensive real-life applications. Secondly, to solve SPM-MM, we design our first algorithm, *Threshold-Adapt*, based on an idea of one popular solution of BMP, *Threshold. Threshold-Adapt* is not scalable for large datasets due to its high time/space complexity. Therefore, we develop another novel algorithm, *Swap-Chain*, which runs faster than *Threshold-Adapt* by orders of magnitude and is scalable to very large datasets (in millions). Finally, we conducted extensive empirical studies on these two solutions.

The rest of this chapter is organized as follows. Section 3.2 defines the SPM-MM problem, and Section 3.3 provides the related work of SPM-MM. Section 3.4 and Section 3.5 introduce two algorithms, *Threshold-Adapt* and *Swap-Chain*, respectively. Section 3.6 gives some discussions. Section 3.7 includes the empirical studies and Section 3.8 concludes the chapter.

## 3.2 The SPM-MM Problem

Let P be a set of service-providers and O be a set of customers. Each service-provider p (customer o) has a service capacity (demand), denoted by p.w (o.w). We represent the Euclidean distance between o and p with d(o, p).

Let  $W_O = \sum_{o \in O} o.w$  and  $W_P = \sum_{p \in P} p.w$ . We assume that the service demands of all customers in O can be satisfied by the service-providers in P, i.e.  $W_P \geq W_O$ . Under this assumption, it is possible that some service-providers are not matched with customers. In case that  $W_P < W_O$ , we swap the roles of P and O and thus this assumption still holds.

**Problem 1 (SPM-MM)** SPM-MM generates the assignment A denoting a set con-

taining the elements in the form of triplets (o, p, w), where (o, p, w) is called a match between o and p and denotes that p provides the service with the amount of w to o. Furthermore, the following three conditions hold.

- No service-provider provides its service of the amount greater than its capacity, i.e.,  $\forall p \in P, \sum_{(o,p,w)\in A} w \leq p.w.$
- Each customer's service demand is satisfied, i.e.,  $\forall o \in O, \sum_{(o,p,w)} w = o.w.$
- The mmd of A is minimized, i.e.,  $\max\{d(o,p)|(o,p,w)\in A\}$  is minimized.

Note that in the following, for clarity, the match (o, p, w) is simply denoted as (o, p) when w = 1.

In order to ease our discussion, we say that an assignment is *full* if it satisfies the Capacity Constraint and the Demand Constraint defined above. Note that there are an exponential number of full assignments. To illustrate, consider the case where |P| = |O| = n and the capacity (demand) of each service-provider (customer) is equal to 1. In this case, there exist n! possible full assignments.

## 3.3 Related Work

We classify the related work into three branches.

**The BMP Problem:** The first branch is the *Bottleneck Matching* problem [32, 8, 25, 28] (BMP). BMP was first proposed by Gross in [32]. Given two sets of n objects,  $A = \{a_1, a_2, ..., a_n\}$  and  $B = \{b_1, b_2, ..., b_n\}$ , and the cost matrix  $C_{n \times n}$  ( $c_{ij}$  represents the cost of matching  $a_i$  with  $b_j$  for  $1 \le i, j \le n$ ), BMP is to find the perfect matching between A and B, which minimizes the maximum matching cost.

One may come up with the following straightforward solution to solve our SPM-MM problem by using the existing solutions for BMP. Specifically, we duplicate each o

in O o.w times and each p in P p.w times. Then, we can use the existing algorithm originally designed for BMP to find the solution for our SPM-MM problem. However, this duplication is cumbersome and undesirable (especially when the capacities/demands are very large), because the resulting datasets would be prohibitively large.

Next, we describe the most popular solution for BMP. [8] provides a comprehensive study of the solutions of BMP, among which, the *Threshold* method has the lowest time complexity. The best-known algorithm for BMP is due to Gabow and Tarjan in [28], which is based on *Threshold*. *Threshold* is based on the property that the minimized maximum matching cost (i.e., the optimal mmd) must reside in the cost matrix  $C_{n \times n}$ . Therefore, it maintains a set X containing the candidates of the optimal mmd, which is initialized to be  $\emptyset$ . For each cost entry c in  $C_{n \times n}$ , it first constructs a bipartite graph between A and B containing the edges each of which is a pair  $(a_i, b_j)$  whose matching cost is at most c. Then, it checks whether there exists a perfect matching in this bipartite graph. If yes, it includes c in X. Finally, it returns the smallest cost in X, which is shown to be the optimal mmd. The above checking operation could be accomplished with a maximum cardinality matching procedure [36] on the corresponding bipartite graph which finds the greatest number of matches in the graph. However, *Threshold* incurs an expensive space cost of  $O(n^2)$  since it has to maintain the cost matrix  $C_{n \times n}$ . Thus, it is not scalable to large datasets.

There is an existing study [25] for BMP in the context of spatial databases where the matching distance between two objects is their Euclidean distance. The method in [25] is exactly the *Threshold* algorithm except that the *maximum cardinality matching* procedure [36] is improved. However, this method cannot be used directly for SPM-MM where the capacities/demands are any positive integers. Besides, the techniques in [25] originally designed for improving the maximum cardinality matching procedure in *Threshold* cannot be adopted for our *Threshold-Adapt* algorithm (which

will be introduced in Section 3.4) since *Threshold-Adapt* involves no maximum cardinality matching procedure.

A *monochromatic* version of BMP (i.e., only one set of data) is considered in [14, 26]. But, these studies are different from ours which uses a *bichromatic* setting where two sets of data (i.e., *P* and *O*) are considered for matching.

Some recent papers [23, 69] in the field of operations research also studied the bottleneck problem and its variations, but they do not focus on the efficiency issue. Specifically, a common technique in this field [23, 69] is constrained optimization/programming, which is known to be slow for large datasets. Besides, [23, 69] only studied the problems in the context of graphs instead of spatial databases.

**Spatial Matching Problems:** The second branch is the existing spatial matching problems [66, 62, 61]. [66] proposed the *SPatial Matching* problem (SPM), which generates a *fair* assignment between *P* and *O*. [62] proposed the *Capacity Constrained Assignment* problem (CCA), which returns the *globally optimized* assignment. Recently, a continuous version of CCA [61] was proposed where customers move *dynamically*.

Since SPM and CCA have different optimization criteria from SPM-MM, the existing solutions developed for SPM and CCA cannot be applied here. In fact, as will be verified in our empirical study, the mmd's of the assignments of SPM and CCA are much larger than the mmd of the SPM-MM assignment.

**Problems with Minimum Maximum Distance:** The third branch is related to some other problems [35, 5] using the *minimum maximum distance* as a measurement. Given n cities, the k-center problem [35], one of the traditional computer science problems, is to build k warehouses at different cities ( $k \le n$ ) such that the maximum distance from a city to its *nearest* warehouse is minimized. The goal of k-center which is to *select* k cities out of n cities is different from that of SPM-MM which is to *match* service-providers and customers. [5] studied an assignment problem between servers

and clients. The matching distance between a server and a client depends on both the physical distance and the *load* of the server where the load of a server corresponds to the number of clients served by this server. In other words, the matching distance between a server and a client defined in [5] in an assignment can be different from the one in another assignment.

# 3.4 Algorithm Threshold-Adapt

### 3.4.1 Theoretical Properties

Given a set P of service-providers and a set O of customers, let  $d_o$  be the optimal mmd for the SPM-MM problem. Intrinsically,  $d_o$  is a pairwise distance between a service-provider p in P and a customer o in O. It follows that  $d_o \in S$ , where S is the set of all possible pairwise distances between P and O, i.e.,  $S = \{d(p,o)|p \in P, o \in O\}$ . Note that  $|S| = |P| \cdot |O|$ . We present this property in the following Lemma 3.4.1.

**Lemma 3.4.1 (Search Space)** Let  $d_o$  be the optimal mmd for the SPM-MM problem.  $d_o$  is in S.

**Proof.**  $d_o$  is a pairwise distance  $(d_o \in S)$ .

According to Lemma 3.4.1, one straightforward method of finding  $d_o$  is to determine whether each value in S is *feasible* for the SPM-MM problem, insert all feasible values into a set X, and find the minimum value in X as  $d_o$ . The definition of "feasibility" is defined next.

**Definition 3.4.1** (Feasibility) Given a positive real number d, d is feasible if and only if there exists a full assignment A between P and O such that its mmd is at most d.

**Lemma 3.4.2 (Feasibility)** Let  $d_o$  be the optimal mmd for the SPM-MM problem.  $d_o$  is feasible.

**Proof.** In the SPM-MM assignment which is full, all matching distances are at most  $d_o$ .

### 3.4.2 Algorithm

We develop our Threshold-Adapt algorithm by using the search space S and the feasibility property described in Lemma 3.4.2. Specifically, Threshold-Adapt checks the feasibility of each distance in S and returns the smallest feasible distance.

**Theorem 3.4.1** The Threshold-Adapt algorithm returns the optimal assignment for the SPM-MM problem.

**Proof.** The correctness follows from Lemma 3.4.1 and 3.4.2.

Let  $\alpha$  be the cost of checking the feasibility of a given value d. One straightforward implementation of *Threshold-Adapt* has the time complexity equal to  $O(|S| \cdot \alpha) = O(|P| \cdot |O| \cdot \alpha)$ . In the following, we consider two issues of *Threshold-Adapt*.

The first issue is to further reduce the size of the search space from  $|P| \cdot |O|$  to  $O(\log(\max\{|P|,|O|\}))$  based on the following *monotonicity* property.

**Lemma 3.4.3** (Monotonicity) Let d and d' be two positive real numbers where d < d'. If d is feasible, then d' is feasible.

**Proof.** Since d is feasible, there exists a full assignment A such that A's mmd is at most d and thus A's mmd is smaller than d'. Therefore, d' is feasible.

According to the above lemma, if we know that a value d' in S is not feasible, then any value d in S smaller than d' must not be feasible. This gives hints for a further reduction of the search space.

Specifically, we sort all values in S in ascending order and store the sorted values in a list L. Then, we adopt binary search to find the smallest feasible value in L (which corresponds to the optimal mmd). This method checks  $O(\log |L|)$  pairwise distances in S. Note that  $|L| = |P| \cdot |O|$ . Thus, the size of the search space becomes  $O(\log(|P| \cdot |O|)) = O(\log(\max\{|P|, |O|\}))$  which is significantly smaller than the original size of  $|P| \cdot |O|$ .

The second issue is to propose an efficient method to perform the judging task to determine whether a given value d is feasible or not. We propose the following three-step algorithm.

Step 1: Construction of a Flow Network wrt d. We construct a flow network  $G_d(V_d, E_d)$  wrt d as follows. We create a source vertex s and a sink vertex t, and  $V_d$  is constructed to be  $P \cup O \cup \{s,t\}$ . For each pair  $(o,p) \in O \times P$  with  $d(o,p) \leq d$ , we create an edge (p,o) in  $E_d$  and set its capacity to be  $\min\{p.w,o.w\}$ . For each p in P (o in O), we create an edge (s,p) ((o,t)) in  $E_d$  and set its capacity to be p.w (o.w).

Step 2: Construction of a Maximum-Flowed Network. We perform a maximum-flow algorithm [3], denoted by  $A_{max-flow}$ , on the flow network  $G_d$  and obtain the maximum flow from s to t in  $G_d$ . We denote the amount of this maximum flow by mf. The maximum-flowed network is the flow network  $G_d$ , where each edge is associated with its flow in the resulting maximum flow. We denote by e.f the flow associated with the edge e.

Step 3: Feasibility Checking on d. We compare mf with  $W_O$ . If  $mf = W_O$ , we conclude that d is feasible; otherwise, we conclude that d is not feasible. In the former case, we construct an assignment, denoted by  $A_d$ , based on the maximum-flowed network at Step 2. We initialize  $A_d$  to  $\emptyset$ . Then, for each edge e in the form of (p, o) in the maximum-flowed network with  $e \cdot f > 0$ , we create a match  $(o, p, e \cdot f)$  in  $A_d$ .

The correctness of the above three-step algorithm is verified by the following lemma.

**Lemma 3.4.4** The three-step algorithm returns a full assignment  $A_d$  with its mmd at most d if and only if d is feasible.

**Proof.** We consider two cases. Case 1:  $mf = W_O$ . In this case, the three-step algorithm returns  $A_d$ . We prove that  $A_d$  is a full assignment with its mmd at most d. First, we know that  $A_d$  satisfies the Capacity Constraint since for each  $p \in P$ ,

$$\sum_{(o,p,w) \in A_d} w = \sum_{e = (p,o) \in E} e.f = (s,p).f \le p.w$$

Second, we show that  $A_d$  satisfies the Demand Constraint by contradiction. Assume there exists a customer  $o' \in O$  whose demand is not satisfied in  $A_d$ , i.e.,  $\sum_{e=(p,o')\in E} e.f < o'.w.$  We have

$$mf = \sum_{e=(p,o)\in E} e.f = \sum_{o''\in O} \sum_{e=(p,o'')\in E} e.f$$

$$< \sum_{o''\in O} o''.w = W_O$$

which contradicts  $mf = W_O$ . Third, it is easy to verify that  $A_d$ 's mmd is at most d since for each edge in the form of (p,o) in E, we have  $d(p,o) \leq d$  (this is guaranteed by Step 1 of the three-step algorithm). In conclusion, we know that  $A_d$  is a full assignment with its mmd at most d which further implies that d is feasible. Case 2:  $mf < W_O$ . In this case, it could be verified that there exists no full assignment which has its mmd at most d by contradiction (note that a full assignment implies a flow with its amount equal to  $W_O$  which contradicts  $mf < W_O$ ). That is, d is not feasible.

**Time Complexity.** After we address the first issue and the second issue, we know that the *Threshold-Adapt* algorithm triggers  $O(\log(\max\{|P|, |O|\}))$  times of run-

ning the maximum-flow algorithm. Thus, the time complexity of *Threshold-Adapt* is  $O(\log(\max\{|P|,|O|\}) \cdot \alpha)$  where  $\alpha$  is the cost of a maximum-flow algorithm (e.g.,  $\alpha = O(n^2m)$  on a flow network with n vertices and m edges if the recently proposed *IBFS* algorithm [31] is adopted). We will test different maximum-flow algorithms in our experiments for optimizing the performance of *Threshold-Adapt*.

We note here that Threshold-Adapt suffers from two intrinsic space problems which limit the application scope of Threshold-Adapt to small/medium-sized datasets only. First, it relies on a search space S whose size is  $|P|\cdot|O|$ . This is prohibitively large when the datasets are large (e.g., S simply occupies about 7.45GB space when |O|=100k and |P|=10k). Second, it has to maintain a flow network  $G_d(V_d,E_d)$  which has its worst-case space complexity of  $O(|P|\cdot|O|)$ . Motivated by the above space issues of Threshold-Adapt, we design another algorithm called Swap-Chain in the next section, which not only avoids these issues by adopting a fundamentally different idea, but also runs faster by orders of magnitude.

## 3.5 Algorithm Swap-Chain

In Section 3.5.1, we give an overview of the *Swap-Chain* algorithm. We then present it in Section 3.5.2, and discuss some issues of *Swap-Chain* and its theoretical results in Section 3.5.3.

#### 3.5.1 Overview

Swap-Chain has the following three steps.

• Step 1 (Assignment Initialization): It first initializes a full assignment A using a given strategy. We will discuss different strategies for this step in Section 3.5.3. One strategy is finding a fair assignment (which is full) by an existing

algorithm [66].

- Step 2 (Assignment Adjustment): It re-assigns some matches in A to form another full assignment A' such that the mmd of A' is smaller than that of A.
- Step 3 (Iterative Step): It repeats Step 2 until it is not possible to perform the assignment adjustment step.

In Step 2, the algorithm reduces the *mmd* of an assignment A by re-assigning some matches in the assignment. Note that the *mmd* of an assignment denotes the maximum matching distance of a match in the assignment and this match is called an *extreme match*. Specifically, the main idea of Step 2 is to find an extreme match in the assignment, break this match and *some* other matches, and re-assign these matches such that the *mmd* of the resulting assignment is smaller.

## 3.5.2 Algorithm

## **Concepts and Algorithm**

Before introducing the *Swap-Chain* algorithm, we introduce some concepts and lemmas related to the algorithm.

Let A be an assignment. Given a customer  $o \in O$ , the *deficient demand* of o in A is defined to be  $o.w - \sum_{(o,p,w)\in A} w.$  o is said to have his/her deficient demand in A if the deficient demand of o in A is non-zero. Otherwise, o is said to have no deficient demand in A. Given a service-provider  $p \in P$ , the *free capacity* of p in A is defined to be  $p.w - \sum_{(o,p,w)\in A} w$ . Similarly, p is said to have its free capacity or have no free capacity in A according to different cases. A service-provider p is said to be *available* in A if it has its free capacity in A. Otherwise, it is said to be *occupied* in A.

**Definition 3.5.1** (d-Available/Occupied service-provider) Given a non-negative real number d and a customer o, a service-provider  $p \in P$  is said to be a d-available

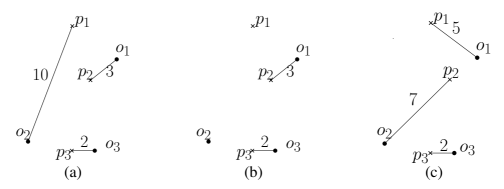


Figure 3.3: The process of Swap-Chain, SPM-MM

service-provider (d-occupied service-provider) for o in A if and only if p is available (occupied) in A and d(o,p) < d.

Example 1 [d-Available/Occupied service-provider] Consider Figure 3.3(b). For the ease of illustration, we assume that the capacity (demand) of each service-provider (customer) is 1 in the figure. Suppose that we have a (non-full) assignment A equal to  $\{(o_1, p_2), (o_3, p_3)\}$ .  $p_1$  is an available service-provider in A but both  $p_2$  and  $p_3$  are occupied service-providers in A. Let d=10. Since  $d(o_1, p_1)=5 < d$ ,  $p_1$  is a d-available service-provider for  $o_1$  in A. Besides, since  $d(o_2, p_2)=7 < d$  and  $d(o_2, p_3)=4 < d$ , both  $p_2$  and  $p_3$  are two d-occupied service-providers for  $o_2$  in A. However, since  $d(o_2, p_1)=10$  which is exactly equal to d,  $p_1$  is not a d-available service-provider for  $o_2$  in A. Note that there does not exist any d-available service-provider for  $o_2$  in a.

**Definition 3.5.2** (d-satisfiability) Given a non-negative real number d and a customer o, o is said to be d-satisfiable in A if and only if one of the following conditions is satisfied.

- Availability Condition: There exists a d-available service-provider for o in A, or
- Non-Availability Condition: There does not exist any d-available serviceprovider for o in A and there exists a d-occupied service-provider p' for o in

A such that p' is matched with another customer o' in A and o' is d-satisfiable in A. In this case, (p', o') is said to be a d-substitute pair for o in A.

Note that "d-satisfiability" is a recursive definition. The availability condition corresponds to the base condition in the recursive definition while the non-availability condition corresponds to the recursive condition.

**Example 2** [d-satisfiability] Consider Example 1. Suppose that the assignment A is still  $\{(o_1, p_2), (o_3, p_3)\}$ . Let d = 10.  $o_1$  is d-satisfiable since there exists a d-available service-provider for  $o_1$  in A (i.e.,  $p_1$ ).  $o_2$  is also d-satisfiable because there does not exist any d-available service-provider for  $o_2$  in A and there exists a d-occupied service-provider for  $o_2$  in A, namely  $p_2$ , such that  $p_2$  is matched with another customer  $o_1$  and  $o_1$  is d-satisfiable in A. Thus,  $(p_2, o_1)$  is a d-substitute pair for  $o_2$  in A.

The following lemma shows the relationship between "d-satisfiability" and the optimal assignment for SPM-MM.

**Lemma 3.5.1 (Optimal Assignment)** Let A be an assignment. If there does not exist any extreme match m in A such that the customer originally matched in m is d-satisfiable in  $A - \{m\}$  where d is the matching distance of m in A, then A is the optimal assignment for the SPM-MM problem.

#### Proof.

We consider two cases. The first case is that for each p and each o, p.w = 1 and o.w = 1. Thus, each match (o', p', w') can be expressed as (o', p') (since w' = 1). The second case is a general case that for each p and each o, p.w and o.w can be equal to any positive integer.

Consider the first case. We prove by contradiction. Let A be the assignment such that there does not exist any extreme match m in A such that the customer o originally

matched in m is d-satisfiable in  $A - \{m\}$  and d is the matching distance of m in A. Suppose that A is not the optimal assignment for the SPM-MM problem. That is, there exists another assignment  $A_o$  such that the mmd of  $A_o$ , denoted by  $d_o$ , is smaller than the mmd of A, denoted by d. Let  $A' = A - \{m\}$ .

Consider  $A_o$ . We know that for each match  $(o', p') \in A_o$ ,  $d(o', p') \leq d_o$ . Since  $d_o < d$ , we have the following. For each match  $(o', p') \in A_o$ ,

$$d(o', p') < d \tag{3.1}$$

There exists a service-provider  $p_1$  such that  $(o, p_1) \in A_o$ . We conclude that

$$d(o, p_1) < d \tag{3.2}$$

Consider A'. We know that o (from match m) is not d-satisfiable in A'. Thus, we deduce that there does not exist any d-available service provider for o in A'. We further consider two sub-cases.

Case (a): There does not exist any d-occupied service-provider for o in A'. We deduce that there does not exist any service-provider p such that d(o, p) < d. This contradicts to that  $d(o, p_1) < d$  (in Inequality (3.2)).

Case (b): There exists a d-occupied service-provider for o in A'. According to Inequality (3.2), we deduce that  $p_1$  is a d-occupied service-provider for o in A'. Thus, there exists a customer  $o_1$  which is matched with  $p_1$  in A'.

In the following, we will show that  $o_1$  is d-satisfiable in A'. After we obtain this result, we can conclude that o is d-satisfiable in A', which leads to a contradiction.

We first construct an undirected graph G (which will be used later in the proof) as follows. Firstly, we construct an assignment  $A_c = (A_o \cup A') - (A_o \cap A')$ . As a result, all customers in O - o are involved in either zero matches in  $A_c$  or *exactly* two distinct matches in  $A_c$ . We construct a set V of vertices to be  $P \cup O$ . For each  $(o', p') \in A_c$ , we

create an edge (o', p'). All edges created form a set E. The graph G is defined based on V and E.

It is easy to verify that in this graph G, any path starting from o is a list containing interleaved customers and service-providers in the form of  $(o, p_1, o_1, p_2, o_2, ...)$  such that the following three rules hold: (R1) o is matched with  $p_1$  in  $A_o$ , (R2)  $o_i$  is matched with  $p_{i+1}$  in  $A_o$  for i = 1, 2, ..., and (R3)  $p_i$  is matched with  $o_i$  in A' for i = 1, 2, ...

According to the three rules, we deduce the following two statements: (1) any path from o to a service-provider in G is non-cyclic, and (2) there exists a path from o to a service-provider point/vertice  $p_n$  such that  $p_n$  is the first service-provider with its free capacity in A' along the path. The correctness of Statement (1) can be shown since there is only one edge involving o in E and each vertice in  $V - \{o\}$  is involved at most two edges in E. Statement (2) can be proved as follows. Since the total number of vertices is bounded by |V| and any path  $\mathcal{P}$  from o to a service-provider is non-cyclic (by Statement (1)), the length of  $\mathcal{P}$  is bounded. Consider a customer o' (not o) along the path  $\mathcal{P}$  from o. Since o' is involved in exactly two edges in E (it is not possible that o' is involved in zero edges in E since o' is along  $\mathcal{P}$  from o), we know that o' is matched in both  $A_o$  and A', and thus the path from o can be prolonged at o'. Consider a service-provider p' along the path  $\mathcal{P}$  from o. If p' is involved in exactly two edges in E, similarly, it is matched in both  $A_o$  and A', and thus the path from o can be prolonged at p'. If it is involved in exactly one edge in E, it means that it is matched in  $A_o$  only (but not A') (by R2) and thus the path from o cannot be prolonged at p'. In this case, p'has its free capacity in A'. This completes the proof when we set  $p_n = p'$  in this case.

Based on the above two statements, we conclude that the path is of the non-cyclic form of  $(o, p_1, o_1, p_2, o_2, ..., p_{n-1}, o_{n-1}, p_n)$  where  $p_n$  is the first service-provider with its free capacity in A' along the path.

Next, we prove that  $o_i$  is d-satisfiable in A' for i = 1, 2, ..., n - 1. We prove by

induction starting from proving the d-satisfiability of  $o_{n-1}$  as a base case. This proof can be done easily by the three rules described above and Inequality (3.1).

Consider  $o_{n-1}$ . Since  $o_{n-1}$  is matched with  $p_n$  in  $A_o$  (by R2), according to Inequality (3.1), we deduce that  $d(o_{n-1}, p_n) < d$ . Thus, since  $p_n$  has its free capacity in A',  $p_n$  is a d-available service-provider for  $o_{n-1}$  in A'. We conclude that  $o_{n-1}$  is d-satisfiable in A'.

Next, we assume that there exists a positive integer k such that  $2 \le k \le n-1$  and  $o_k$  is d-satisfiable in A'. With this assumption, we want to prove that  $o_{k-1}$  is d-satisfiable in A'. Since  $p_n$  is the first service-provider with its free capacity in A' along the path, we know that  $p_k$  has no free capacity in A' (since  $0 \le k \le n-1$ ). Besides,  $0 \le k \le n-1$ ). Besides,  $0 \le k \le n-1$ . According to Inequality (3.1), we have  $0 \le k \le n-1$ . We deduce the following Statement (\*):  $0 \le k \le n-1$ .

Consider two cases. Case (I): There exists a d-available service-provider for  $o_{k-1}$  in A'. In this case,  $o_{k-1}$  is d-satisfiable in A' (by Availability Condition).

Case (II): There does not exist any d-available service-provider for  $o_{k-1}$  in A'. In this case, note that  $p_k$  is matched with  $o_k$  in A' (by R3) and  $o_k$  is d-satisfiable in A' (by the hypothesis). Since  $p_k$  is a d-occupied service-provider for  $o_{k-1}$  in A' (by Statement (\*)), we deduce that  $o_{k-1}$  is d-satisfiable in A' (by Non-Availability Condition).

In Case (I) and Case (II), we have the same conclusion. By induction, we conclude that  $o_i$  is d-satisfiable in A' for i=1,2,..,n-1.

So,  $o_1$  is d-satisfiable in A'. The proof for Case 1 is complete.

Consider Case 2. Case 2 can be easily transformed to Case 1 by duplicating each customer o' (service-provider p') o'.w (p'.w) times, and each match (o', p', w') w' times. This completes the proof.

## **Algorithm 5** Algorithm Swap-Chain(P, O)

- 1: initialize a full assignment A between P and O
- 2: **while** there exists an extreme match m in A which involves a customer o such that o is d-satisfiable in  $A \{m\}$  where d is the matching distance of this extreme match  $\mathbf{do}$
- 3:  $A \leftarrow Swap(A, m)$
- 4: return A

The above lemma motivates us to design Swap-Chain as shown in Algorithm 5. In this algorithm, Swap is the re-matching operation related to an extreme match m in A. We will describe how we perform this operation next.

# The Swap Operation

We first need to introduce a concept called "d-swapping chain" which is used for the Swap operation. Roughly speaking, it is a list of objects describing which customers and service-providers in the current assignment are involved in the re-matching (or Swap) operation such that the new matching distance for each of these customers is smaller than d where d is a non-negative real number.

A list is represented in the form of  $(x_1, x_2, ..., x_l)$  where  $x_i$  is an object (either a customer or a service-provider) for  $i \in [1, l]$  and l is the number of objects in the list. Given a list L in the form of  $(x_1, x_2, ..., x_l)$ , a pair in the form of  $(x_i, x_{i+1})$  is said to be an even pair in L if i is divisible by 2. Otherwise, it is said to be an odd pair in L. Given two lists  $L_1$  and  $L_2$  where  $L_1$  is  $(x_1, x_2, ..., x_l)$  and  $L_2$  is  $(y_1, y_2, ..., y_{l'})$ , the list concatenation of  $L_1$  and  $L_2$ , denoted by  $L_1 \diamond L_2$ , is defined to be  $(x_1, x_2, ..., x_l, y_1, y_2, ..., y_{l'})$ .

**Definition 3.5.3** (d-Swapping Chain) Let A be an assignment. Suppose that o is d-satisfiable in A. We define a d-swapping chain from o in A, denoted by  $C_d(o)$ , as follows according to the availability condition and the non-availability condition.

•  $C_d(o)$  is the list (o, p') if the availability condition is satisfied where p' is a d-available service-provider for o in A, or

•  $C_d(o)$  is the list  $(o, p') \diamond C_d(o')$  if the non-availability condition is satisfied where (p', o') is a d-substitute pair for o in A.

*Example 3* [d-Swapping Chain] Consider Example 1. The assignment A is still  $\{(o_1,p_2),(o_3,p_3)\}$ . Let d=10. A d-swapping chain from  $o_1$  in A, denoted by  $C_d(o_1)$ , can be  $(o_1,p_1)$ . Besides, a d-swapping chain from  $o_2$  in A, denoted by  $C_d(o_2)$ , can be  $(o_2,p_2)\diamond C_d(o_1)$  (which is equal to  $(o_2,p_2,o_1,p_1)$ ) since  $(p_2,o_1)$  is a d-substitute pair for  $o_2$  in A.

Let A be an assignment and d be a non-negative real number at least the mmd of A. Given a customer o, a d-swapping chain from o in A, denoted by C, has the following properties.

- The total number of objects in C is even.
- C is a list containing interleaved customers and service-providers. The first object in C is a customer. We call it as the first customer wrt C. The last object in C is a service-provider p' and the second-to-last object in C is a customer o'. We call p' as the last service-provider wrt C. Note that p' is a d-available service-provider for o' in A.
- Each *odd* pair in L is in the form of (o', p') and d(o', p') < d.
- Each *even* pair in L is in the form of (p', o') and  $d(p', o') \leq d$  (note that d(p', o') is at most the *mmd* while d is at least the *mmd* as has been specified). For each even pair (p', o') in L, there exists a positive integer w' such that  $(o', p', w') \in A$  and w' is said to be the *weight* of the even pair (p', o').

Note that given a customer o and a non-negative real number d at least the mmd of an assignment A, o is d-satisfiable in A if and only if there exists a d-swapping chain from o in A.

After we describe the d-swapping chain, we are ready to describe how to perform the re-matching operation, Swap, based on this chain. For the ease of illustration, we first assume that the capacity (demand) of each service-provider (customer) is 1. We call this assumption the *unit assumption*. After we explain the intuition under the unit assumption, we will relax it. Under the unit assumption, the amount of the service given by a service-provider to a customer in a match is exactly equal to 1. Thus, the weight of each possible even pair in a swapping chain C is equal to 1.

Suppose that we are given a full assignment A. We describe our Swap algorithm as follows.

**Step (a) (Extreme Match Breaking):** We find an extreme match m. Let d be the matching distance of m in A and o be the customer matched in m. We then break this extreme match m in A. That is, we remove m from A and form a new assignment A' (i.e.,  $A' = A - \{m\}$ ).

**Step (b) (Swapping Chain Finding):** We then find a d-swapping chain from o in A', denoted by C.

**Step (c) (Chain Breaking):** Note that each *even* pair (p', o') in C corresponds to a match in A'. For each even pair (p', o') in C, we break the match (p', o') (or formally (o', p')) in A'. Note that the customer o' in each even pair (p', o') in C has no deficient demand before this step but has his/her deficient demand after this step.

**Step (d) (Chain Matching):** For each *odd* pair (o', p') in C, we form a match (o', p') in A'. At this moment, the customer o' in each odd pair (o', p') in C has no deficient demand.

Let X be the set of customers involved in the swapping chain C. Note that with the above Swap algorithm, the mmd, say d', of the resulting assignment involving only the customers in X is smaller than the mmd, say d, of the original assignment involving only the customers in X. This is because we make sure that for each odd pair (o', p')

in C (which forms a match in the resulting assignment), the distance between o' and p' is smaller than d.

If the original assignment contains  $exactly \ one$  extreme match, it is easy to see that the mmd of the resulting assignment involving all customers is smaller than the mmd of the original assignment involving all customers. However, it is possible that multiple extreme matches exist in an assignment A which have the same matching distance d. The mmd of the resulting assignment involving all customers decreases only after we break all of these extreme matches.

Example 4 [Swap] Suppose that the capacity (demand) of each service-provider (customer) is 1. Consider Figure 3.3(a) which shows a full assignment  $\{(o_1, p_2), (o_2, p_1), (o_3, p_3)\}$ . We denote this assignment by A.  $(o_2, p_1)$  is an extreme match in A. Let  $d = d(o_2, p_1) = 10$ . The Swap operation based on A and match  $(o_2, p_1)$  works as follows. First, we break the extreme match  $(o_2, p_1)$  and the resulting assignment is shown in Figure 3.3(b). Second, we find a d-swapping chain C from  $o_2$  which is  $(o_2, p_2, o_1, p_1)$  (Refer Example 3 for illustration). Third, we break the even pairs in C which include  $(p_2, o_1)$  only. Forth, for each odd pair in C, we form its corresponding match and thus matches  $(o_2, p_2)$  and  $(o_1, p_1)$  are formed. Figure 3.3(c) shows the resulting assignment. Clearly, the new assignment is still full, but with a smaller mmd (i.e., 7).

Next, we relax the unit assumption such that the capacity (demand) of each service-provider (customer) could be any positive integer instead of 1. In this case, the weight of an even pair in a swapping chain C can be different from that of another even pair.

The *Swap* algorithm can also be used with this relaxation except the following changes related to the weight of a match.

Step (a) (Extreme Match Breaking): We perform the same operation as before. But,

after the breaking of an extreme match in the form of (o, p, w), resulting an assignment A', we obtain that o has its deficient demand equal to w (instead of 1) while p has its free capacity at least w (instead of 1).

**Step (b) (Swapping Chain Finding):** Similarly, we perform the same operation.

Step (c) (Chain Breaking): In this step, due to the weights of matches, we have to calculate the weights of matches which are used in this chain breaking operation. Specifically, let W be  $W_e(C) \cup \{w_o\} \cup \{w_p\}$  where  $W_e(C)$  is the set of the weights of all possible even pairs in C,  $w_o$  is the deficient demand of the first customer wrt C in A' and  $w_p$  is the free capacity of the last service-provider wrt C in A'. We define the *swapping amount* of the chain C, denoted by Amount(C), to be  $\min_{w \in W} \{w\}$ . Roughly speaking, Amount(C) corresponds to the greatest possible amount of service in a match along the chain such that Steps (c) and (d) can be executed successfully. Let  $w_s = Amount(C)$ . Note that  $w_s$  is smaller than or equal to the weight of each even pair in C.

We execute Step (c) as follows. For each *even* pair (p', o') in C, we break the match (o', p', w') in A' where w' is a positive integer and form a match  $(o', p', w' - w_s)$  in A'. Note that the customer o' in each even pair (p', o') in C has no deficient demand before this step but has his/her deficient demand equal to  $w_s$  after this step.

**Step (d) (Chain Matching):** In Step (d), we perform the matching with the weight  $w_s$ . That is, for each *odd* pair (o', p') in C, we form a match  $(o', p', w_s)$  in A'. Note that at this moment, the customer o' in each odd pair (o', p') in C except the first odd pair has no deficient demand in A'.

Step (e) (Deficient Demand Checking): This step is new. If the customer o in the first odd pair has no deficient demand in A' (this case happens when  $w_s = w_o$ ), we can return the resulting assignment A' generated from Step (d). If o has his/her deficient demand in A' (this case occurs when  $w_s < w_o$ ), then we continue to execute Step (b)

## **Algorithm 6** Algorithm Swap(A, m)

```
Require: a full assignment A and an extreme match m in A
 1: // Step (a) (Extreme Match Breaking)
 2: Let m be the match involving customer o and service-provider p with its matching
    distance equal to d
 3: A' \leftarrow A - \{m\}
 4: while there exists a d-swapping chain from o in A' do
      // Step (b) (Swapping Chain Finding)
       C \leftarrow a d-swapping chain from o in A'
 6:
      // Step (c) (Chain Breaking)
 7:
       w_s \leftarrow Amount(C)
 8:
       for each even pair (p', o') in C do
 9:
         find a match (o', p', w') in A'
10:
          A' \leftarrow A' - \{(o', p', w')\}
11:
         if w' \neq w_s then A' \leftarrow A' \cup \{(o', p', w' - w_s)\}
12:
       // Step (d) (Chain Matching)
13:
      for each odd pair (o', p') in C do
14:
          A' \leftarrow A' \cup \{(o', p', w_s)\}
15:
       // Step (e) (Deficient Demand Checking)
16:
       if o has no deficient demand in A' then break;
17:
18: // Step (f) (Post-Matching)
19: if o has his/her deficient demand in A' equal to w'' then
       A' \leftarrow A' \cup \{(o, p, w'')\}
21: return A'
```

to Step (d) until o has no deficient demand in A' or o becomes not d-satisfiable in A'. When we stop the above iterative process, if o has no deficient demand in A', similarly, we can return A' as the output. If o is not d-satisfiable in A', we will run an additional step called *Post-Matching* in Step (f).

**Step (f) (Post-Matching):** This step is also new. It will be executed if o has his/her deficient demand in A', say w'', and is not d-satisfiable in A'. In this case, it is not possible to reduce the matching distance of the extreme match m involving o. Thus, we create the match (o, p, w'') in A' where p is the service-provider involved in m. Finally, we return A'.

The pseudo-code of *Swap* is shown in Algorithm 6.

With Algorithm 6, it is easy to show the correctness of the *Swap-Chain* algorithm (Algorithm 5) as follows.

**Theorem 3.5.1** The Swap-Chain algorithm returns the optimal assignment for the

SPM-MM problem.

**Proof.** The correctness follows from Lemma 3.5.1.

## 3.5.3 Remaining Issues & Theoretical Analysis

**Remaining Issues**. There are two remaining issues in *Swap-Chain*, namely the initialization of a full assignment (line 1 in Algorithm 5) and the Swapping Chain Finding step in the *Swap* algorithm (line 6 in Algorithm 6).

Issue 1: There are many possible ways of initializing a full assignment. In our implementation, we consider the following two methods, namely Sort and Fair. Sort returns an assignment by a two-step approach. First, for each  $o \in O$ , it maintains a list of all service-providers in ascending order of their distances to o. Second, it processes all  $o \in O$  one by one. When processing a specific o, it traverses the service-providers in o's corresponding list sequentially and for the currently traversed p, it assigns the service with the amount equal to  $\min\{o.d, p.f\}$  from p to o, where o.d is o's deficient demand and p.f is p's free capacity. The traversing process stops when o's demand has been satisfied. Fair denotes the method of generating the fair assignment. Note that we do not adopt the globally optimized assignment in the initialization since the time complexity of the algorithm for finding the globally optimized assignment is much higher than that for finding other assignments like the fair assignment.

Issue 2: For Swapping Chain Finding (i.e., finding a d-swapping chain from a customer o in an assignment A), we design a Breadth First Search (BFS) method as follows. It maintains a queue Q and initially inserts o into Q. Then, it processes the elements in Q one by one as follows. It starts processing the first element in Q. If the current element in Q (being processed) is a customer, say  $o_c$ , it inserts into Q all service-providers (in any order) that have their distances from  $o_c$  smaller than d and have not been inserted into Q. These service-providers can be found by issuing a

range query on P from  $o_c$ . We say that  $o_c$  is the *parent* of all these service-providers. If the current element in Q (being processed) is a service-provider, say  $p_c$ , consider two cases. Case 1:  $p_c$  has no free capacity. In this case, it inserts all the customers matched with  $p_c$  in A into Q (in any order) and  $p_c$  is said to be the *parent* of all these customers. Case 2:  $p_c$  has its free capacity. In this case, it traces all *ancestors* of  $p_c$  until the (starting) customer o is reached, and returns the traced list (in this list, the first element is o and the last element is  $p_c$ ) as a d-swapping chain from o. The above process continues with the next element in Q until either a d-swapping chain is found or all elements in Q have been processed. In the latter case, it means that there does not exist any d-swapping chain from o.

Here, we need to perform range queries on P. Let  $\beta(|P|)$  be the cost of a range query on a dataset of size |P|. In [15], with the data structure with its size of  $O(|P|(\log |P| \log \log |P|)^2)$  and its construction time complexity of  $O(|P| \log |P|)$ ,  $\beta(|P|) = O(\log |P| + k)$  where k is the size of the answer of this query. In practice, k << |P| usually holds. Note that in our implementation, instead of the data structure proposed in [15], we adopt an R-tree index built on P for supporting range queries since it is available in commercial databases and is found to be efficient in practice (though it does not have good worst-case asymptotic performance).

**Theoretical Properties**. We first describe some theoretical properties which will be used to analyze the time complexity of our *Swap-Chain* algorithm.

Given a match (o, p, w) in an assignment, we say that (o, p) is its *match signature*. Given an assignment A, a list of interleaved objects from P and O in the form of  $(o_1, p_1, o_2, p_2, \ldots, o_n, p_n)$  is said to be a *match cycle* if each two adjacent objects in the list form a match in A, i.e.  $o_i$  is matched with  $p_i$  for  $1 \le i \le n$ ,  $o_{i+1}$  is matched with  $p_i$  for  $1 \le i \le n - 1$ , and  $o_1$  is matched with  $p_n$  in the assignment. The length of a cycle is defined to be the number of elements in the cycle. An assignment A is said to be cyclic if A contains a match cycle.

Interestingly, a non-cyclic assignment has a theoretical bound on the number of matches in the assignment.

**Lemma 3.5.2** Given P and O, the number of matches in a non-cyclic assignment is bounded by |P| + |O| - 1.

**Proof.** First, we show that in a non-cyclic assignment A involving no match cycle, there exists an element e (either a service-provider or a customer) such that e is involved in *exactly* one match in A (We can prove by contradiction since if each element is involved in at least two matches, there exists a match cycle in the assignment). We say this match is *critical*. Second, given an assignment A, we iteratively remove each *critical* match from A until no matches exist in A. Since each removal operation makes at least one element unmatched and the last removal operation makes exactly two elements unmatched, we know the number of matches in A is at most |P| + |O| - 1.

Furthermore, given an assignment A with a match cycle C, the following lemma suggests that C could be *destroyed* in A easily such that some conditions in A are still satisfied.

**Lemma 3.5.3** Let A be a cyclic assignment with a match cycle C. We can transform A to another assignment A' such that (1) the mmd of A' is at most that of A; (2) the deficient demand (free capacity) of each  $o \in O$   $(p \in P)$  remains unchanged and (3) A' does not contain C nor any matches with new match signatures compared with A. Besides, the cost of this transformation is O(n) where n is the length of C.

**Proof.** Let the cycle C be  $(o_1, p_1, o_2, p_2, ..., o_n, p_n)$ . Without loss of generality, let  $(o_1, p_1)$  be the match along C which has the smallest matching weight, says  $w_m$ . We

break each match  $(o_i, p_i, w_i)$  with the amount of  $w_m$  for each  $i \in [1, n]$ . Thus,  $o_i$  has its deficient demand at least  $w_m$  and  $p_i$  has its free capacity at least  $w_m$  for each  $i \in [1, n]$ . Next, we create a new match  $(o_{i+1}, p_i, w_m)$  for each  $i \in [1, n-1]$  and a new match  $(o_1, p_n, w_m)$ . (Note that for the new match formed  $(o, p, w_m)$ , if there exists an original match (o, p, w) in the assignment, we just combine these two matches as a single match  $(o, p, w_m + w)$ ). Let A' be the resulting assignment. It can be verified that  $o_1$  and  $p_1$  originally matched in A are not matched in A' and thus A' does not contain cycle C. Besides, it is easy to verify that for each  $o \in O$ ,  $\sum_{(o,p,w)\in A} w = \sum_{(o,p,w)\in A'} w$  and for each  $p \in P$ ,  $\sum_{(o,p,w)\in A} w = \sum_{(o,p,w)\in A'} w$ . Furthermore, the mmd of A' is at most that of A and no matches with new match signatures are formed in A'. Clearly, the cost of the above process is simply O(n).

Time Complexity. We let |V| = |P| + |O| and  $|E| = |P| \cdot |O|$ . Suppose that we build an index as introduced in [15] on P to facilitate range queries described before. Let  $\lambda$  be the time complexity of building this index. Let  $\gamma$  be the time complexity of the full assignment initialization (line 1 of Algorithm 5). Let R be the total number of possible extreme matches fetched in Swap-Chain (i.e., the number of iterations in lines 2-3 of Algorithm 5). Let R denote the time complexity of the R algorithm. The time complexity of R algorithm is R algorithm is R and R algorithm.

Consider  $\lambda$ . From [15], we know that  $\lambda = O(|P| \log |P|)$ .

Consider  $\gamma$ . If *Sort* is adopted, it could be verified that  $\gamma$  is equal to  $O(|O| \cdot |P| \log |P|)$ . If *Fair* is used,  $\gamma$  is equal to  $O((|P| + |O|) \cdot (\log |P| + \log |O|))$  [66]. Besides, we introduce a lemma which will be used later.

*Lemma 3.5.4* The assignment initialized by Sort and the assignment initialized by Fair are both non-cyclic. □

**Proof.** Let  $A_s$  be the assignment initialized by *Sort* and  $A_f$  be the one initialized

by Fair.

We first prove that  $A_s$  is non-cyclic. Suppose Sort processes O in order of  $o_1, o_2, ..., o_m$ , where m is the size of O. We denote by  $A_{o_i}$  the assignment that is formed immediately after processing  $o_i$  (thus,  $A_{o_m} = A_s$ ). We claim that for a specific customer  $o_i$ , among all products that are matched with  $o_i$  in  $A_{o_i}$ , at most one product has its free capacity non-zero. This can be verified by the principle adopted in Sort that  $o_i$  always exhausts the current product chosen to be matched with  $o_i$  before the next product is considered. Now, we show  $A_s$  is non-cyclic by contradiction. Assume that there exists in  $A_s$  a match cycle  $C = (o_{c_1}, p_{c_1}, ..., o_{c_n}, p_{c_n})$ . Without loss of generality, among all customers involved in C, we assume  $o_{c_1}$  is the first customer processed by Sort. Consider  $A_{oc_1}$  (the assignment formed immediately after  $o_{c_1}$  is processed). Both  $p_{c_1}$  and  $p_{c_n}$  are matched with  $o_{c_1}$  in  $A_{oc_1}$  (since  $p_{c_1}$  and  $p_{c_n}$  are matched with  $o_{c_1}$  in  $A_{oc_1}$  (since  $p_{c_1}$  and  $p_{c_n}$  are matched with  $oc_1$  in  $oc_2$  ( $oc_2$ ) later on where processing  $oc_2$  ( $oc_2$ ). Thus, this leads a contradiction that at most one product matched with  $oc_1$  in  $oc_2$  has its free capacity non-zero.

Next, we prove that  $A_f$  is non-cyclic by contradiction. This proof can be done by using the fact that no dangling pair [66] exists in a fair assignment. We only consider the case where the pairwise distances between P and O are distinct. For the other case where some pairwise distances are identical, we can always perform an infinitesimal perturbation on the locations of some products and/or customers such that ties of distances are broken. Assume that there exists a match cycle  $C = (o_1, p_1, ..., o_n, p_n)$  in  $A_f$ . Without loss of generality, we assume that match  $(o_1, p_1, w_1)$  has the smallest matching distance among all matches involved in C. As a result, we know  $d(o_1, p_1) < d(o_2, p_1)$  and  $d(o_1, p_1) < d(o_1, p_n)$ , which implies that  $(o_1, p_1)$  is a dangling pair [66]. However, according to [66], there exist no dangling pairs for a fair assignment. There, we know  $A_f$  is not a fair assignment, which leads to a contradic-

tion.

Consider R. Before we give the bound on R, we give a lemma.

**Lemma 3.5.5** A match with a given match signature can be fetched as an extreme match at most once in Swap-Chain.

**Proof.** This lemma is trivially true if all pairwise distances are distinct. This lemma also holds even if they are not distinct. This is because during the execution of *Swap* (in line 3 of *Swap-Chain*), once the extreme match with a particular match signature is broken, no matches with the same match signature will be formed again except the last Step (f) (i.e., post-processing) which denotes that there is no need to fetch additional extreme matches and the algorithm terminates.

Note that there are at most |E| (=  $|P| \cdot |O|$ ) possible match signatures. By Lemma 3.5.5, we deduce that R is bounded by |E|. In practice, R << |E|. In our experiments, R is about 500 on average, which is very small compared with |E| which is as large as 250,000,000 in our default setting.

Consider I. According to Algorithm 6, I depends on the cost of the while-loop (lines 6-17) and the total number of while-loops, denoted by t. Consider a while-loop which involves the operation of finding a d-swapping chain (line 6), whose cost is denoted by  $\mathcal{C}_1$ , the operation of re-matching the elements along the chain (line 7-17), whose cost is denoted by  $\mathcal{C}_2$ , and an additional operation introduced here which is used to transform the assignment obtained to a non-cyclic assignment and whose cost is denoted by  $\mathcal{C}_3$ . Thus, I is  $t \cdot (\mathcal{C}_1 + \mathcal{C}_2 + \mathcal{C}_3)$ .

Consider  $C_1$  which corresponds to the time cost of the BFS implementation. Note that at the beginning of each while-loop, the assignment is non-cyclic. This is because the initialized assignment is non-cyclic (Lemma 3.5.4) and at the end of each while-loop, the additional operation introduced here transforms the assignment to a non-

cyclic one. Thus, it could be verified that  $C_1 = O(|O| \cdot \beta(|P|) + |P|)$  (the BFS method (1) involves at most |O| range queries on P (which incurs the cost of  $O(|O| \cdot \beta(|P|))$ ), and (2) retrieves at most |P| + |O| - 1 matches (from service-providers) according to Lemma 3.5.2 and the fact that the assignment is non-cyclic (which incurs the cost of O(|P| + |O|))).

Consider  $C_2$ . It is simply O(|P| + |O|) (=O(|V|)).

Consider  $\mathcal{C}_3$ . After the Chain Matching step, the assignment contains O(|V|)matches (since it contains at most |P| + |O| - 1 matches at the beginning of the whileloop and the Chain Matching step forms at most  $\min\{|P|, |O|\}$  new matches). Clearly, each match cycle in an assignment A corresponds to a cycle in the un-directed graph  $G_A(V', E')$ , which involves P and O as vertices in V' and all matches as edges in E'. Note that |E'| = O(|V|) (by Lemma 3.5.2). Thus, to find a match cycle in A, we can find a cycle in  $G_A$  and this can be easily achieved by a common DFS technique [21], which runs in O(|V'| + |E'|) (=O(|V|)) time. According to Lemma 3.5.3, destroying a match cycle incurs O(|V|) (a match cycle has its length at most |P| + |O|) and it does not introduce any match with a new match signature. So, we can transform the assignment to a non-cyclic one by iteratively destroying the match cycles until no match cycles exist in the assignment. Thus,  $C_3 = O(c \cdot |V|)$ , where c is the number of match cycles formed due to the Chain Matching step. It could be verified that c is bounded by  $\min\{|P|, |O|\}$  since the Chain Matching step introduces at most  $\min\{|P|, |O|\}$  matches and each such match can form at most one new match cycle. In practice,  $c << \min\{|P|, |O|\}$  (e.g., c is about 17 on average in our experiments under the default setting).

Consider t. Recall that t is the number of while-loops in Swap needed to re-satisfy the deficient demand of customer o due to the break operation on the extreme match involving o. Clearly, t is bounded by  $\overline{w} = \min\{\max_{p \in P} p.w, \max_{o \in O} o.w\}$ . Usually,

t is much smaller than this upper bound  $\overline{w}$ . For example, in our experiments on real datasets, on average, t is 2 (with a maximum of 40) but  $\overline{w}$  is in thousands.

In view of the above discussion, we know that  $I = O(t \cdot (|V| \cdot \beta(|P|) + c \cdot |V|))$ . As a result, the time complexity of Swap-Chain is  $O(\lambda + \gamma + R \cdot t \cdot (|V| \cdot \beta(|P|) + c \cdot |V|))$ , where R << |E|,  $t << \min\{\max_{p \in P} p.w, \max_{o \in O} o.w\}$ , and  $c << \min\{|P|, |O|\}$ .

# 3.6 Discussion

Any assignment with its *mmd* equal to the optimal *mmd* is a solution of SPM-MM. Thus, there may exist multiple possible solutions for SPM-MM. In this case, our Swap-Chain returns one of them at random. However, SPM-MM can be enriched by considering a secondary objective (e.g., minimizing the sum of the matching distances) for the final solution among these multiple solutions. Furthermore, the bottleneck nature of the SPM-MM objective makes it quite easy to be incorporated with a secondary objective since the optimized mmd, say  $d_o$ , can always be used as a hard constraint for optimizing the secondary objective. Specifically, matching any pair of two objects which has its distance bounded by  $d_o$  does not destroy the optimality while matching any pair of two objects which has its distance larger than  $d_o$  definitely ruins the optimality. Thus, we can adopt a two-step mechanism for the SPM-MM problem with a secondary objective. First, we compute the optimal mmd, say  $d_o$ , using Swap-Chain. Second, we ignore all pairs (o, p) with  $d(o, p) > d_o$  for matching when optimizing the secondary objective. For instance, if the secondary objective is to minimize the sum of the matching distances, we can solve this enriched version of SPM-MM easily by first computing the optimal mmd  $d_o$  and then adopting any popular algorithm for MinimumWeight Matching [3] with the constraint that all pairs (o, p) with  $d(o, p) > d_o$  cannot be matched (this could be achieved by excluding from the graph used by the algorithm all those edges with the corresponding distances larger than  $d_o$ ).

Next, we discuss SPM-MM in a more general setting where the pairwise distances between P and O could be non-metric or non-spatial. Interestingly, our proposed methods can also be adapted to this general setting. Threshold-Adapt still works in the general setting (recall that *Threshold-Adapt* is adapted from *Threshold* which is designed for general bipartite graphs). We can also adapt Swap-Chain to the general setting with some sacrifice of its time complexity as follows. Two parts involved in Swap-Chain rely on the spatial setting, namely the Fair method for initializing a full assignment (a fair one) and the BFS method for finding a d-swapping chain in an assignment A. To initialize a fair assignment in the general setting, one can adopt the Stable Marriage algorithm which incurs the cost of  $O(|P| \cdot |O|)$  [29] (instead of  $O((|P| + |O|) \cdot (\log |P| + \log |O|))$  in the spatial setting [66]). To find a d-swapping chain from a customer o in the general setting, one can first materialize a directed graph G'(V', E') such that (1)  $V' = P \cup O$ ; (2) for each  $o \in O$ , (o, p) is a directed edge in E' for all  $p \in P$  with d(o, p) < d; (3) for each  $p \in P$ , (p, o) is a directed edge in E' for all  $o \in O$  that is matched with p in A, and then find a path from o to a service-provider p with its free capacity non-zero using a BFS on G'. The resulting path corresponds to a d-swapping chain from o in A. Clearly, |V'| = |V| and  $|E'| \le |E|$ . Thus, the cost of the BFS method is O(|V| + |E|) (instead of  $O(|O| \cdot \beta(|P|) + |P|)$  in the spatial setting, where  $\beta(|P|)$  is the range query cost on P).

Finally, we would like to note some differences between our d-swapping chain technique and the well-known  $augmenting\ path$  techniques. A typical augmenting path technique is used for computing the  $maximum\ flow$  whose main idea is to iteratively finding an  $augmenting\ path$  and augmenting the flow along this path until no augmenting paths are possible. As could be noticed, the goal of an augmenting path technique is to increase the flow iteratively while the goal of our d-swapping chain technique is to keep the flow while decreasing the mmd of the corresponding matching. Specifically, in our d-swapping chain technique, we find an extreme match (o, p) and break it so

	Cardinality		
	Populated	Fire Sta-	
	Areas (PA)	tions (FS)	
AB	4,999	447	
BC	6,609	595	
ON	12,474	1,215	
QC	12,936	1,259	

Table 3.1:	Real	datasets	(for	SP-
M-MM)			,	

Factor	Configuration		
Cardinality $( O )$	10k, 30k, <b>50k</b> , 70k, 100k		
Dim.	2, 3, 4, 5		
Size ratio $(r)$	5, <b>10</b> , 15, 20, 25		
Weight ratio (k)	1, 1.5, <b>2</b> , 2.5, 3		
O's weights	[1, 10)		
Scalability $( O )$	250k, 500k, 750k, 1000k		

Table 3.2: Synthetic datasets (for SPM-MM)

that we will find a d-swapping chain from o, while in an augmenting path technique, there is no such breaking operation on a chosen match before the augmenting path is to be found.

# 3.7 Empirical studies

We used four real datasets, namely AB, BC, ON and QC, in our experiments. Each real dataset contains two sets of spatial objects, a set of populated areas (PA) and a set of fire stations (FS). Specifically, dataset AB contains the set of PAs and the set of FSs in Alberta, Canada. Datasets BC, ON and QC contain the same information in the three other provinces in Canada, namely, British Columbia, Ontario and Quebec, respectively. We collected the PAs from Census Canada (http://www12.statcan.gc.ca), each of which corresponds to a dissemination area, and estimated the coordinates of PAs with the help of the Postal Code Conversion File of Canada [60]. The population of each PA ranges from 400 to 700 in most cases [60]. We collected the FSs from Fire-Canada (http://www.firecanada.ca) and estimated the coordinates of FSs via Google Maps. The capacities of FSs range from 5,500 to 10,000. The coordinates are all normalized to range [0,10000]. For each dataset, we adopt the set of PAs as *O* and the set of FSs as *P*. The summaries of the real datasets are shown in Table 3.1.

We also used synthetic datasets in our experiments, which are generated as follows. The coordinates of spatial objects follow the Uniform distribution on range [0, 10000] by default. The demand of each customer in O is set to be [1, 10) randomly. To generate the capacities of the service-providers in P, we define a parameter k, called weight ratio, to be the expected ratio between the sum of the service capacities of all service-providers and the sum of the service demands of all customers, i.e.,  $k = \sum_{p \in P} p.w / \sum_{o \in O} o.w$ . Based on the configuration of k, we set the capacities of the service-providers. By default, the capacities are set to be [80, 120) randomly. The parameter configuration of synthetic datasets is shown in Table 3.2 where the default values are shown in bold font.

## 3.7.1 SPM-MM vs. Existing Spatial Matching

We conducted experiments to compare the optimal mmd,  $mmd_o$ , with the mmd's of the fair assignment and the globally optimized assignment, namely  $mmd_{fair}$  and  $mmd_{global}$ , respectively. In this experiment, we randomly select 10% (5%) in P and 10% (5%) in Q for each real (synthetic) dataset. This is because the algorithm (we use the SSPA algorithm in [3]) for computing  $mmd_{global}$  is not scalable to large datasets.

Figure 3.4 shows that  $mmd_{fair}$  and  $mmd_{global}$  are larger than  $mmd_o$ . For example, in the real dataset ON (Figure 3.4(a)), the ratio between  $mmd_{fair}$  ( $mmd_{global}$ ) and  $mmd_o$  is about 3.5 (2.3). We have similar results on synthetic datasets as shown in Figure 3.4(b).

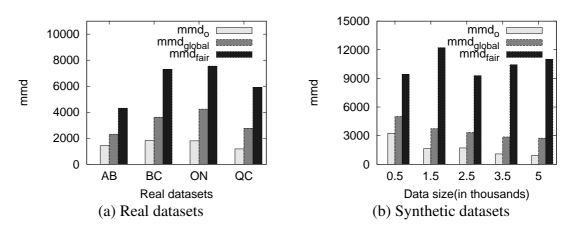


Figure 3.4: Results for the *mmd*'s of different assignments, SPM-MM

## 3.7.2 Performance Study

Next, we give the performance study on our proposed algorithms, namely *Threshold-Adapt* and *Swap-Chain*, which include eight instances in total. The details are described as follows.

Recall that *Threshold-Adapt* involves a *maximum-flow* procedure. In the literature, many maximum-flow algorithms have been developed which could be categorized into three branches, namely, *Augmenting-Path* (which mainly includes Dinic, BK and IBFS [31]), *Push-Relabel* (which mainly includes HIPR and PRF), and *Pseudoflow* (which mainly includes HPR). More details about these maximum-flow algorithms could be found in [64] (and the references therein). Besides, according to [64], these maximum-flow algorithms usually favor different applications and it is not always the case that a maximum-flow algorithm with a smaller time complexity runs faster than another with a larger one. Motivated by this, we consider all the above six maximum-flow algorithms, namely, Dinic, BK, IBFS, HIPR, PRF and HPF, for optimizing *Threshold-Adapt*, and the corresponding instances of *Threshold-Adapt* are denoted by *TA-Dinic*, *TA-BK*, *TA-IBFS*, *TA-HIPR*, *TA-PRF* and *TA-HPF*, respectively. Besides, we consider two instances of the *Swap-Chain* algorithm, namely, *Swap-Fair* and *Swap-Sort*, with the initialization methods of *Fair* and *Sort*, respectively.

We evaluated the algorithms mainly in terms of *running time* and *memory*, and study the effects of cardinality, dimensionality, size ratio and weight ratio on the performance of the algorithms. The memory of *Threshold-Adapt* is mainly due to the search space S and the flow network graph, and the memory of *Swap-Chain* is mainly due to the R-tree built on P and the maintained assignment.

We implemented our algorithms in C/C++ and conducted the experiments on a Linux platform with a 2.26GHz CPU and 36GB physical memory.

We present our experimental results as follows.

(1) Effect of Cardinality. We vary |O| and the results are shown in Figure 3.5. We have the following observations. First, there is a clear efficiency gap between the Swap-Chain algorithms and the Threshold-Adapt algorithms and the gap becomes larger when the data size increases. For example, when |O| = 100k, Swap-Chain is faster than TA-IBFS by more than one order of magnitude. Second, the two Swap-Chain algorithms favor different cases. Specifically, Swap-Sort runs faster than Swap-Fair on relatively small datasets (e.g.,  $\leq 40k$ ) while the opposite case becomes true on relatively large datasets. This could be explained by the fact that (1) Swap-Sort has no cost of building an R-tree on O while Swap-Fair does and (2) Swap-Sort has a more expensive initialization procedure (i.e., Sort) than Swap-Fair. Third, the memory usages of the Swap-Chain algorithms are quite low while those of the Threshold-Adapt algorithms are dramatically higher (by 2-3 orders of magnitude). For example, when |O| = 100k, the Swap-Chain algorithms use less than 50MB while each of Threshold-Adapt algorithms occupies more than 15GB memory. Forth, among all Threshold-Adapt algorithms, TA-IBFS runs the fastest and occupies the least memory. For ease of presentation, in the following, we focus on TA-IBFS only as the representative of the Threshold-Adapt algorithms since it beats all other instances of Threshold-Adapt in terms of both time efficiency and space efficiency.

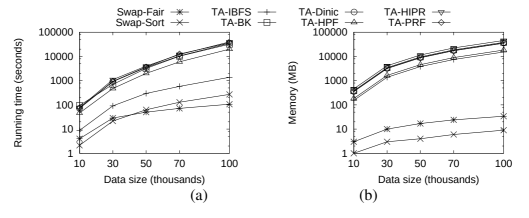


Figure 3.5: Effect of cardinality (synthetic datasets, SPM-MM)

(2) Effect of Dimensionality. Figure 3.6 shows the results of the effect of dimen-

sionality. We observe that the dimensionality only affects the Swap-Fair algorithm slightly. Specifically, when the dimensionality increases, the running time of Swap-Fair increases slightly. This is because Swap-Fair needs to build the R-trees on both P (for searching d-swapping chains) and O (for computing a fair assignment), which cost increases when the dimensionality increases. The dimensionality has negligible effects on Swap-Sort and TA-IBFS.

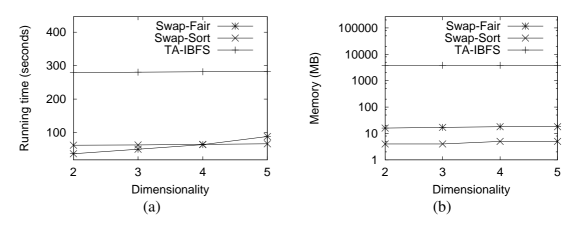


Figure 3.6: Effect of dimensionality (synthetic datasets, SPM-MM)

(3) Effect of Size ratio. We observe some opposite trends on running time and memory when we increase the size ratio r compared with those when we increase the data size. This is reasonable since when the size ratio r increases, |P| decreases (note that |O| is fixed).

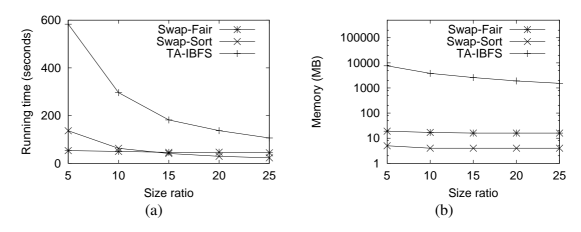


Figure 3.7: Effect of size ratio (synthetic datasets, SPM-MM)

(4) Effect of Weight ratio. Figure 3.8 shows the effect of the weight ratio k. We

observe that the weight ratio has slight effect on TA-IBFS only. Specifically, when k increases, the running time of TA-IBFS decreases slightly. The reason might be that when k increases (i.e., the total capacities of the service-providers becomes relatively larger), it is more likely that an augmenting path (note that IBFS is an augmenting path algorithm) carries more flow and thus the process of computing the maximum-flow could be finished more quickly.

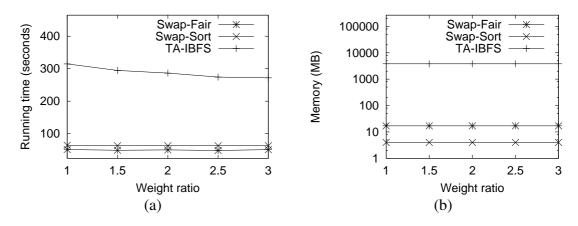


Figure 3.8: Effect of weight ratio k (synthetic datasets, SPM-MM)

- (5) Scalability test. Figure 3.9 shows the results of the scalability test for *Swap-Sort* and *Swap-Fair*. Since *Threshold-Adapt* is not scalable, we did not conduct this test for *Threshold-Adapt*. As shown in the figure, the two algorithms are still efficient on large datasets (in millions). Furthermore, *Swap-Fair* is more scalable than *Swap-Sort*. This is because on a large dataset, the initialization process of *Swap-Fair* (i.e., *Fair*) is much faster than that of *Swap-Sort* (i.e., *Sort*).
- (6) Experiments on real datasets. Figure 3.10 shows the results for real datasets which are similar to the results for synthetic datasets. Compared with the *Threshold-Adapt* algorithm, our *Swap-Chain* algorithms run faster and use significantly less memory.
- (7) Comparison with the *Threashold* algorithm in Euclidean space. We are interested in studying the performance of our proposed algorithms when they are used for

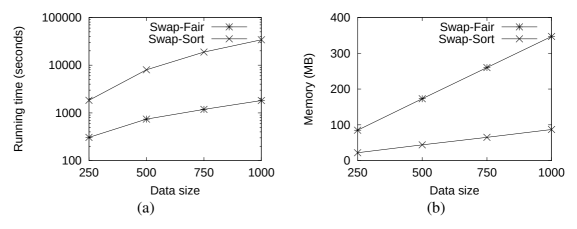


Figure 3.9: Scalability test (synthetic datasets, SPM-MM)

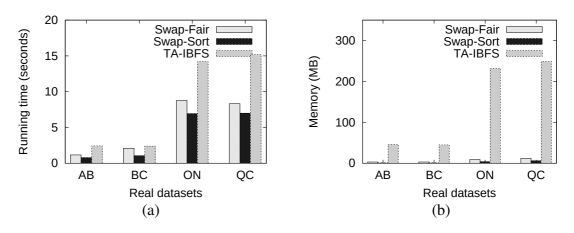


Figure 3.10: Results for real dataset, SPM-MM

the un-weighted version of SPM-MM (i.e., all the capacities/demands are 1's). We compare our algorithm with the state-of-the-art called Match [25] which has a theoretical time complexity of  $O(n^{1.5} \log n)$ . We note here that though Match has a smaller time complexity, it has quite a narrow application scope (i.e., for the un-weighted version only) and the time complexity is restricted to the 2D space only [25]. The results are shown in Figure 3.11. We observe that our Swap-Chain algorithms have comparable running time with the Match algorithm and run even faster than Match on relatively large datasets. This might be due to the fact that a constant factor which could be large is omitted from the time complexity analysis in [25]. Besides, we found that our Swap-Chain algorithms enjoy the superiority of space efficiency over the Match algorithm. We also used our real datasets for this experiment by setting the capacities/demands to 1s and observed similar results.

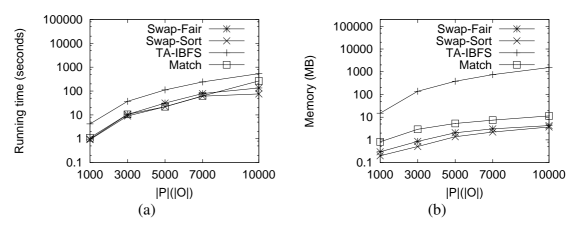
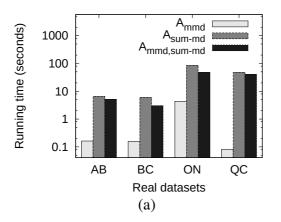


Figure 3.11: Threshold vs. Swap-Chain, SPM-MM

(8) Experiments with a Secondary Objective. Besides, we conducted some experiments on the SPM-MM problem with a secondary objective of minimizing the *sum of matching distances* called *sum-md*. Let  $A_{mmd}(A_{sum-md})$  be the assignment obtained by optimizing mmd (sum-md) only. Let  $A_{mmd,sum-md}$  be the assignment obtained by optimizing mmd first and sum-md second. We adopted the SSPA algorithm [3] for optimizing sum-md. We conducted our experiments on both synthetic and real datasets where each synthetic/real dataset was sampled first with the sampling rate set to 5% due to the relatively expensive cost of SSPA. The results on the real datasets are shown in Figure 3.12. We observe that on average, compared to  $A_{sum-md}$ ,  $A_{mmd,sum-md}$  can be obtained with a similar time (the cost of optimizing sum is an additional part but the constraint of the optimized sum helps reduce the running time of the process of optimizing sum and has the sum-sum value usually not far away from the sum-sum value of sum-sum (e.g., within 1.1 factor).

(9) Experiments with non-Euclidean Distances. In addition, we conducted some experiments on our proposed algorithms (*Threshold-Adapt* and *Swap-Chain*) when they are applied to the cases where non-Euclidean distances are used. We used the same real datasets except that the underlying distances between pairs of two objects are measured by the driving time between the two objects. The results are shown in Figure 3.13. We observe that compared with the case of using the Euclidean distances, the efficiency



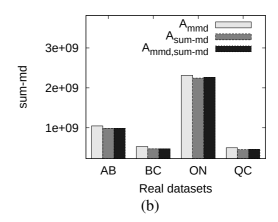
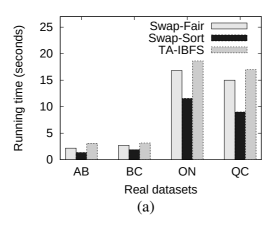


Figure 3.12: Experiments with a secondary objective, SPM-MM

of *Threshold-Adapt* is similar while the efficiency of the *Swap-Chain* algorithms, especially *Swap-Fair*, degrades to some extent. But, the *Swap-Chain* algorithms still retain the superiority over *Threshold-Adapt* in terms of both time efficiency and space efficiency. For example, on dataset QC, the running time of *Swap-Fair* (*Swap-Sort*) is about 15s (9s) while that of *Threshold-Adapt* is nearly 17s. Thus, compared with the spatial setting, the degrading ratio of *Swap-Fair* (*Swap-Sort*) is about 9/15 (8/9) and that of *Threshold-Adapt* is about 16/17.



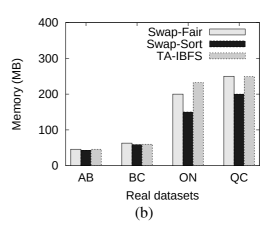


Figure 3.13: Experiments with non-Euclidean distances, SPM-MM

(10) Comparison with the Augmenting Path Technique. We also conducted experiments on the *Swap-Chain* algorithms with the adaptions of <u>Augmenting Path</u> (AP) techniques [3]. We interpret our *d*-swapping chains as augmenting paths and denote the resulting *Swap-Chain* algorithms corresponding to *Swap-Fair* and *Swap-Sort* by

AP-Fair and AP-Sort, respectively. In our implementations of AP-Fair and AP-Sort, when finding an augmenting path which corresponds to finding a d-swapping chain in Swap-Fair and Swap-Sort, respectively, we do a BFS in a graph structure G which contains the edges (o,p) for all pairs of (o,p) with d(o,p) < d and also the edges (p,o) for all matches (o,p,w) in the current assignment. Figure 3.14 shows the results. We observe that there is a clear efficiency gap between Swap-Fair (Swap-Sort) and AP-Fair (AP-Sort), and this gap becomes larger when the data size increases. Besides, AP-Fair (AP-Sort) occupies significantly more memory than Swap-Fair (Swap-Sort). The reason for the efficiency gap is that each range query on P in Swap-Fair (Swap-Sort) is performed in  $O(\log |P| + k)$  time [15] where k is the size of the answer of the query while its counterpart in AP-Fair (AP-Sort) is performed by scanning an adjacent list which takes O(|P|) time. The reason for the results of memory usage is that Swap-Fair (Swap-Fair (Swap

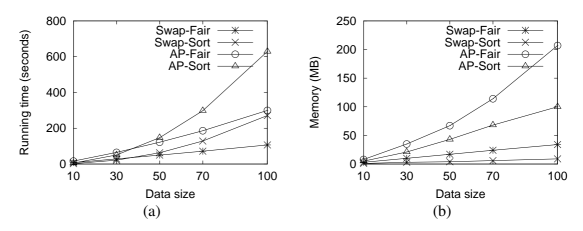


Figure 3.14: d-swapping chain vs. Augmenting path, SPM-MM

**Conclusion:** The *Swap-Chain* algorithms, which are efficient and scalable, beat the *Threshold-Adapt* algorithms in terms of both time efficiency and space efficiency. Besides, *Swap-Sort* runs faster than *Swap-Fair* when the datset is relatively small (e.g.,  $|O| \le 40k$ ) while *Swap-Fair* enjoys its superiority over *Swap-Sort* on large datasets.

# 3.8 Conclusion

In this chapter, we propose a new problem called <u>SPatial Matching for Minimizing Maximum matching distance</u> (SPM-MM). We design two algorithms for SPM-MM, namely <u>Threshold-Adapt</u> and <u>Swap-Chain</u>. <u>Threshold-Adapt</u> is simple and easy to understand but not scalable to large datasets. <u>Swap-Chain</u> avoids the scalability issues of <u>Threshold-Adapt</u> by adopting a novel idea of swapping the matches iteratively and runs faster than <u>Threshold-Adapt</u> by orders of magnitudes. We conducted extensive experiments which verified the efficiency and scalability of <u>Swap-Chain</u>.

# CHAPTER 4

# DIRECTION-PRESERVING TRAJECTORY SIMPLIFICATION

## 4.1 Introduction

With the proliferation of GPS-embedded devices (e.g., smart phones and taxis), trajectory data is becoming ubiquitous. Indeed, it has been studied extensively in the past decades in the literature of Moving Objects Databases (MOD) [59, 53]. Trajectory data is usually generated by periodically collecting the position of a moving object with the help of the GPS technologies.

Since the raw trajectory data is usually very large, simplifying trajectory data is important. To appreciate this, consider a city with 10k taxis. Suppose that we track the trajectory of each taxi by sampling its position once every 5 seconds (i.e., the sampling rate is 5s). The size of the collected trajectories for just one day is approximately 4 GB.

Raw trajectory data is large, and hence expensive to store. Even worse, it is expensive to manipulate and to analyze on account of its large size. In fact, most existing query processing and data mining algorithms on trajectory data are memory-resident and thus cannot be used with raw trajectory data that is too large to fit in memory.

A question one may ask is why not just sampling less frequently to reduce the size of the data. The answer is that, in real life, objects have great variance in their velocities. A taxi moving at 40 mph would have moved about 100 yards in 5s, whereas another taxi stuck at a traffic signal may not have moved at all. Obviously, we need

more frequent observations of the former than of the latter. Similarly, we need more observations to capture a taxi that makes a turn and fewer for one that continues straight. Therefore, standard practice is to oversample initially, and then to simplify by eliminating observations that add little information.

In view of this, several algorithms have been developed for simplifying trajectory data [50, 24, 55, 51, 40]. All these algorithms make the natural assumption that the goal should be to simplify trajectories such that the *position information* captured in the simplified trajectories is "similar" to the position information captured in the original trajectories. We can call them *position-preserving trajectory simplification algorithms*. However, as we will soon see, this objective, though natural, is not the best choice in many situations. To illustrate, let us work through a toy example in detail.

**Example 5 (Motivating Example)** Consider three raw trajectories  $T_1$ ,  $T_2$  and  $T_3$  as shown in bold lines in Figure 4.1(a)(i), (a)(ii) and (a)(iii), respectively. Each of these trajectories has *four* positions,  $p_1$ ,  $p_2$ ,  $p_3$  and  $p_4$ .  $T_1$  and  $T_2$  are *similar* to each other, and each of them is *dissimilar* to  $T_3$ . Thus, a trajectory clustering algorithm, such as [46], should group  $T_1$  and  $T_2$  in the same cluster and place  $T_3$  by itself in a separate cluster.

Now suppose that these raw trajectories are too large, and so must be simplified to three points each before being further processed. We could use an existing position-preserving trajectory simplification, denoted by  $\mathcal{A}_{pos}$ , for this simplification. Following existing studies, the first position  $p_1$  and the last position  $p_4$  in each trajectory have to be kept. Therefore, one of position  $p_2$  and position  $p_3$  is to be retained, and the other one dropped.

Consider the simplification process on  $T_1$ . It can drop either  $p_2$  or  $p_3$  in the simplified trajectory. Let  $d_1$  ( $d_2$ ) be  $p_2$ 's ( $p_3$ )'s perpendicular distance to line segment  $\overline{p_1p_3}$  ( $\overline{p_2p_4}$ ). Since  $d_1>d_2$ ,  $\mathcal{A}_{pos}$  drops  $p_3$  and returns the simplified trajectory  $T_1'$  as

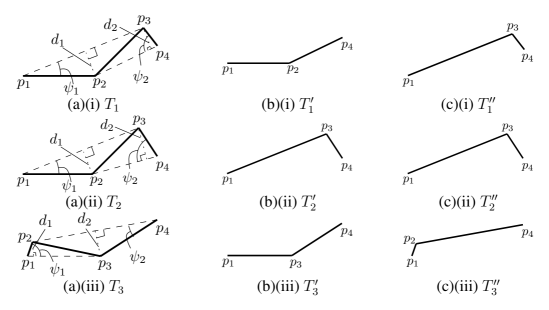


Figure 4.1: A motivating example (for DPTS)

shown in Figure 4.1(b)(i). Similarly,  $\mathcal{A}_{pos}$  return  $T_2'$  (Figure 4.1(b)(ii)) and  $T_3'$  (Figure 4.1(b)(iii)) as the simplified trajectories of  $T_2$  and  $T_3$ , respectively. We now see that, though raw trajectories  $T_1$  and  $T_2$  are *similar*, their simplified trajectories  $T_1'$  and  $T_2'$  generated by  $\mathcal{A}_{pos}$  are *dissimilar*. On the other hand, raw trajectories  $T_1$  and  $T_3$  are *dissimilar*, but their simplified trajectories  $T_1'$  and  $T_3'$  generated by  $\mathcal{A}_{pos}$  are *similar*. In consequence, the clustering algorithm on the simplified trajectories  $T_1'$ ,  $T_2'$  and  $T_3'$ , places  $T_1$  and  $T_3$  together into one cluster, and thus fails to produce correct (or expected) clusters.

In contrast, as will be shown later, a *direction-preserving* trajectory simplification method we introduce below, denoted by  $\mathcal{A}_{dir}$ , would simplify  $T_1$ ,  $T_2$  and  $T_3$  to  $T_1''$  (Figure 4.1(c)(i)),  $T_2''$  (Figure 4.1(c)(ii)) and  $T_3''$  (Figure 4.1(c)(iii)), respectively. Since  $T_1''$  and  $T_2''$  are similar to each other and each of them is dissimilar to  $T_3''$ , the clustering algorithm based on these simplified trajectories would produce the expected clusters.

Before we can discuss direction-preserving trajectory simplification in depth, we first have to describe what direction information is, which we do next.

## **4.1.1 Direction Information**

When an object moves from position p to position p', we define the *direction* of this movement to be the angle of an anticlockwise rotation from the positive x-axis to a vector from p to p'. The directions of all movements captured in the trajectory is called the *direction information*, and is used heavily, both directly and indirectly, in a wide range of applications on trajectory data. We list some of them as follows.

- *Map Matching* [6]. Given a digital map of a road network and a trajectory of an object moving on the road network, the map matching problem is to locate the trajectory on the digital map. Since each road segment in the road network has its own *orientation*, restricting the *directions* of the movements in the trajectory, the direction information plays an essential role in most map matching algorithms [6].
- *Knowledge Discovery on Trajectory Data*. As with other types of data, a rich set of knowledge discovery tasks has been proposed on the trajectory data [30]. Among them, many algorithms rely heavily on the direction information, which include [46, 37] for *Clustering*, [44] for *Outlier Detection* and [45] for *Classification*.
- *Direction-based Query Processing*. Sometimes, there are reasons to query trajectory information directly. One example is to find the trajectories moving within a direction range in a given time slot [7]. Another example is to find trajectories *similar* to a given trajectory, where the similarity measurement is based solely on moving direction [54].

In short, there are many situations in which direction preservation is important. Furthermore, as we show analytically in Section 4.3 and empirically in Section 4.6, direction preservation is stronger than position preservation, in that a simplification that

preserves direction information well can be shown to preserve position information also, within some reasonable bounds. However, the converse is not true: position-preserving simplifications can be very bad at direction preservation.

## **4.1.2** Direction-Preserving Trajectory Simplification (DPTS)

In this thesis, we propose a new trajectory simplification mechanism called *Direction-Preserving Trajectory Simplification* (DPTS) such that the direction information loss due to the simplification process is bounded. Within DPTS, we propose a *direction-based measurement*  $E_d$ , which is new and is defined to measure the *error* of a simplified trajectory in terms of the direction information. Let T be a trajectory and T' be a simplification of T. The *error* (or *simplification error*) of T' under  $E_d$ , denoted by  $\epsilon(T')$ , is equal to the maximum *angular difference* between the direction of the movement during each time period in T and the direction of the movement during the same time period in T'. Then, the problem of DPTS is to simplify a given trajectory such that its *size* is minimized and its incurred simplification error (i.e.,  $\epsilon(T')$ ) is bounded by a given error tolerance  $\epsilon_t$  where  $\epsilon_t \in [0, \pi)$ .

We use the *maximum* angular difference rather than the average angular difference to preserve better the shape of the trajectory. If we used the average one, we could still have a few segments that were completely off, resulting in the types of errors illustrated in Figure 4.1 for position-preserving techniques.

In this thesis, we study the properties of DPTS, develop multiple algorithms to solve the DPTS problem, both exactly and approximately, and evaluate our algorithms experimentally. Specifically, we make the following contributions.

**Contribution.** First, we propose a novel notion of direction-preserving trajectory simplification, which favors a wide spectrum of applications on trajectory data. Second, we show that DPTS not only preserves direction information, but also preserves posi-

tion information, thereby supporting a wide range of applications. Third, we adopt a common dynamic programming (DP) technique for DPTS. Since it is not scalable, we propose a novel optimal algorithm called SP for DPTS. SP solves DPTS by first constructing a graph based on the given trajectory, then computing a shortest path in this graph and finally returning the solution for DPTS according to the shortest path found. The time complexity of SP is  $O(C \cdot n^2)$ , where C is usually a small constant (C=1 if  $\epsilon_t \leq \pi/2$ ). Fourth, since even an  $O(n^2)$  running time is likely to be unacceptable for a large n, we propose a scalable approximate algorithm called *Intersect* which runs in O(n) time. We show that *Intersect* provides a certain degree of the quality guarantee in terms of the size of the simplified trajectory returned, in spite of running so fast. Finally, we perform a careful experimental comparison of these algorithms and a baseline using real trajectory data. The baseline is developed by common sense modifications of standard trajectory simplification techniques to address the DPTS problem.

The rest of this chapter is organized as follows. We define the DPTS problem in Section 4.2 and review the related work in Section 4.3. We introduce the optimal and approximate algorithms of DPTS in Section 4.4 and Section 4.5, respectively. We give the empirical study in Section 4.6 and conclude this chapter in Section 4.7.

## **4.2** Problem Definition

A trajectory is represented by a sequence of n triplets in the form of  $((x_1,y_1,t_1),(x_2,y_2,t_2),...,(x_n,y_n,t_n))$ , where  $(x_i,y_i)$  is the position in the 2D Euclidean space at time stamp  $t_i$ . We define *positions*  $p_i=(x_i,y_i)$  for each  $i\in[1,n]$ . Then, T's trace is the sequence of ordered positions, i.e.,  $(p_1,p_2,...,p_n)$ .

Since the direction information of a trajectory is captured by its trace only, in the following, following existing studies, we focus on the trace part of the trajectory and use the terms "trajectory" and "trace" interchangeably. Thus, we simply denote T by

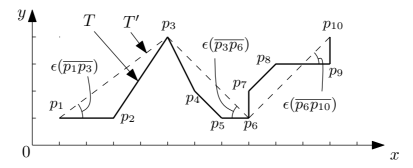


Figure 4.2: A running example (for DPTS)

 $(p_1, p_2, ..., p_n)$  by keeping the position information only. The *size* of T, denoted by |T|, is defined to be the number of positions in T.

Consider a running example as shown in Figure 4.2. In this figure, the trajectory T is represented in the form of  $(p_1, p_2, ..., p_{10})$ . The size of this trajectory (i.e., |T|) is 10. The start position of T is  $p_1$  and the end position of T is  $p_{10}$ .

The straight line linking two positions  $p_i$  and  $p_j$  in T where  $1 \le i < j \le n$  is denoted by  $\overline{p_ip_j}$ . If  $p_i$  and  $p_j$  are *adjacent* in T (i.e., j=i+1), then  $\overline{p_ip_j}$  is said to be a *segment* in T. Thus, a trajectory could also be regarded as a sequence of n-1 segments joining at n-2 positions (in addition to unique start and end positions).

In Figure 4.2, the solid horizontal straight line connecting  $p_1$  and  $p_2$  is denoted by  $\overline{p_1p_2}$ . Similarly, the dashed inclined straight line connecting  $p_1$  and  $p_3$  is denoted by  $\overline{p_1p_3}$ . Here,  $\overline{p_1p_2}$  is a segment in T but  $\overline{p_1p_3}$  is not a segment in T. All segments in T are shown in solid lines in the figure. In T, there are 9 segments jointing at 8 positions, namely  $p_2, p_3, ..., p_9$ .

Trajectory T' is said to be a *simplification* of T if T' is of the form of  $(p_{s_1}, p_{s_2}, ..., p_{s_m})$  where  $m \leq n$  and  $1 = s_1 < s_2 < ... < s_m = n$ . Note that  $p_1$  and  $p_n$  in T must be kept in any simplification of T. There are m-1 segments in T', and T' is using m-1 segments to represent T containing n-1 segments. For each  $k \in [1, m)$ , the segment  $\overline{p_{s_k}p_{s_{k+1}}}$  in T' is used to approximate the sequence of segments between  $p_{s_k}$  and  $p_{s_{k+1}}$  in T, namely  $\overline{p_{s_k}p_{s_{k+1}}}$ ,  $\overline{p_{s_k+1}p_{s_k+2}}$ , ...,  $\overline{p_{s_{k+1}-1}p_{s_{k+1}}}$ . In

other words, this sequence of segments in T is approximated by a single segment (i.e.,  $\overline{p_{s_k}p_{s_{k+1}}}$ ) in T' only.

Consider our running example. Let  $T'=(p_1,p_3,p_6,p_{10})$ . T' is a simplification of T in Figure 4.2. Here,  $s_1=1,s_2=3,s_3=6$  and  $s_4=10$ . Note that the size of T' is 4. All segments in T' are shown in dashed lines in the figure. There are 3 segments in T'. In other words, T' is using 3 segments to approximate 9 segments in T. Consider segment  $\overline{p_1p_3}$  in T'. It is used to approximate the sequence of segments between  $p_1$  and  $p_3$  in T, namely  $\overline{p_1p_2}$  and  $\overline{p_2p_3}$ . In other words,  $\overline{p_1p_2}$  and  $\overline{p_2p_3}$  are approximated by a single segment  $\overline{p_1p_3}$ . Similarly, trajectory  $T''=(p_1,p_{10})$  is also a simplification of T, which uses only one segment (i.e.,  $\overline{p_1p_{10}}$ ) to approximate the whole trajectory T.

Direction-based Error Measurement  $E_d$ . Given a segment  $\overline{p_ip_{i+1}}$  in T, the direction of  $\overline{p_ip_{i+1}}$ , denoted by  $\theta(\overline{p_ip_{i+1}})$ , is defined to be the angle of an anticlockwise rotation from the positive x-axis to a vector from  $p_i$  to  $p_{i+1}$ . Thus, each direction falls in  $[0,2\pi)$ . Consider our running example (Figure 4.2).  $\theta(\overline{p_7p_8})$  is  $\pi/4(=0.788)$  radian and  $\theta(\overline{p_4p_5})$  is  $7\pi/4(=5.498)$  radian, as illustrated in Figure 4.3(a). It is easy to verify that  $\theta(\overline{p_1p_2})$  is equal to 0 radian,  $\theta(\overline{p_2p_3})$  is equal to 0.983 radian (=  $\tan^{-1} 3/2$ ) and  $\theta(\overline{p_1p_3})$  is equal to 0.644 radian (=  $\tan^{-1} 3/4$ ).

The angular difference between two directions  $\theta_1$  and  $\theta_2$ , denoted by  $\triangle(\theta_1, \theta_2)$ , is defined to be the minimum of the angle of the anticlockwise rotation from  $\theta_1$  to  $\theta_2$  and that from  $\theta_2$  to  $\theta_1$ , i.e.,

$$\triangle(\theta_1, \theta_2) = \min\{|\theta_1 - \theta_2|, 2\pi - |\theta_1 - \theta_2|\} \tag{4.1}$$

For illustration, Figure 4.3(b) shows the case where  $\triangle(\theta_1, \theta_2) = |\theta_1 - \theta_2|$  and Figure 4.3(c) shows the case where  $\triangle(\theta_1, \theta_2) = 2\pi - |\theta_1 - \theta_2|$ . Note that the angular difference between any two directions falls in  $[0, \pi]$ .

Consider our running example. The angular difference between  $\theta(\overline{p_1p_2})$  and

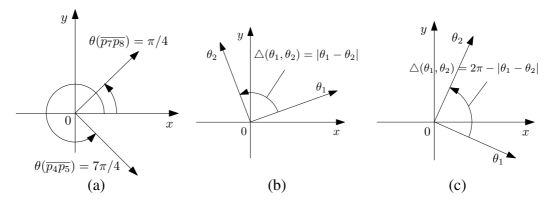


Figure 4.3: Examples illustrating the definition of "direction" and "angular difference", DPTS

 $\theta(\overline{p_1p_3})$  is |0 - 0.644| = 0.644 and that between  $\theta(\overline{p_2p_3})$  and  $\theta(\overline{p_1p_3})$  is |0.983 - 0.644| = 0.339.

Let  $T'=(p_{s_1},p_{s_2},...,p_{s_m})$  be a simplification of T The simplification error of T' under  $E_d$ , denoted by  $\epsilon(T')$ , is defined as follows. Given a segment  $\overline{p_{s_k}p_{s_{k+1}}}$  in T', the simplification error of  $\overline{p_{s_k}p_{s_{k+1}}}$ , denoted by  $\epsilon(\overline{p_{s_k}p_{s_{k+1}}})$ , is defined to be the greatest angular difference between the direction of  $\overline{p_{s_k}p_{s_{k+1}}}$  and the direction of a segment in T approximated by  $\overline{p_{s_k}p_{s_{k+1}}}$ . That is,

$$\epsilon(\overline{p_{s_k}p_{s_{k+1}}}) = \max\nolimits_{s_k \leq h < s_{k+1}} \triangle(\theta(\overline{p_{s_k}p_{s_{k+1}}}), \theta(\overline{p_hp_{h+1}}))$$

Then, the *simplification error* of T' under  $E_d$  is defined to be the *greatest* simplification error of a segment in T'. That is,

$$\epsilon(T') = \max_{1 \le k < m} \epsilon(\overline{p_{s_k} p_{s_{k+1}}}) \tag{4.2}$$

Consider back our running example (Figure 4.2). Each segment in T' has its simplification error. Consider the first segment  $\overline{p_1p_3}$  in T' which approximates two segments in T, namely  $\overline{p_1p_2}$  and  $\overline{p_2p_3}$ . Recall that  $\triangle(\theta(\overline{p_1p_3}), \theta(\overline{p_1p_2})) = 0.644$  and  $\triangle(\theta(\overline{p_1p_3}), \theta(\overline{p_2p_3})) = 0.339$ . Thus, the simplification error of  $\overline{p_1p_3}$  (i.e.,  $\epsilon(\overline{p_1p_3})$ ) is equal to  $\max\{0.644, 0.339\} = 0.644$ . Similarly, we compute the simplification errors of the second segment  $\overline{p_3p_6}$  and the third segment  $\overline{p_6p_{10}}$  in T' which are both

equal to 0.785. Thus, the simplification error of T' in this example is equal to  $\max\{0.644, 0.785, 0.785\} = 0.785$ .

In the following, when we write  $\epsilon(\overline{p_ip_j})$   $(0 \le i < j \le n)$ , we mean the simplification error of  $\overline{p_ip_j}$  when it is used to approximate the line segments between  $p_i$  and  $p_j$  in T.

**Problem Statement of DPTS.** Let T be a trajectory and  $\epsilon_t$  be the error tolerance  $(\epsilon_t < \pi)$ . Trajectory T' is said to be an  $\epsilon_t$ -simplification of T if T' is a simplification of T and  $\epsilon(T') \le \epsilon_t$ .

The DPTS problem is formalized as follows.

**Problem 2 (DPTS)** Given a trajectory T and an error tolerance  $\epsilon_t$ , the DPTS problem is to find the  $\epsilon_t$ -simplification of T with the smallest size.

Consider our running example. Suppose that we set  $\epsilon_t$  to 0.785. T' is an  $\epsilon_t$ -simplification of T since  $\epsilon(T') = 0.785 \le \epsilon_t$ . In fact, T' is the  $\epsilon_t$ -simplification of T with the smallest size (which involves only four remaining positions).

# 4.3 Analysis of Previous Work

We describe how DPTS relates to existing error measurements (Section 4.3.1) and trajectory simplification techniques (Section 4.3.2).

## **4.3.1** Existing Error Measurements

In this section, we show that the direction-preserving simplified trajectories give error guarantees in position-related properties, such as length and speed. However, the reverse is not true. That is, the position-preserving simplified trajectories [50, 51, 55, 40, 24] do not give any error guarantee on the direction information.

Before we give our claims/properties (Section 4.3.1), we review some representative existing error measurements (Section 4.3.1).

#### **Existing Error Measurements**

Let  $T=(p_1,p_2,...,p_n)$  be a trajectory and  $T'=(p_{s_1},p_{s_2},...,p_{s_m})$  be a simplification of T ( $m \le n$ ). Several *position-based* measurements for evaluating the "simplification error" of T' have been defined in the literature. These measurements for T' are usually defined to be a *distance measure* which takes T and T' as input. For each position  $p_h$  of T at the time stamp equal to  $t_h$  where  $1 \le h \le n$ , the distance measure defines the *estimated position* of  $p_h$ , denoted by  $p'_h$ , in T' based on some criteria. Let  $d(\cdot,\cdot)$  be the Euclidean distance between two given points. Thus, the distance measure is defined to be  $\max_{h \in [1,n]} d(p_h,p'_h)$ . Since different distance measures have different definitions on estimated positions, in the following, we focus on how to define estimated positions for some representative distance measures.

- (1) Closest Euclidean Distance: With this distance measure,  $p'_h$  is defined to be the location on the segment  $\overline{p_{s_k}p_{s_{k+1}}}$  of T' with  $s_k \leq h \leq s_{k+1}$ , which has the smallest Euclidean distance from  $p_h$ . We define a mapping function  $M_{\mathcal{C}}$  which maps  $p_h$  and T' to  $p'_h$  for this distance measure. That is,  $p'_h = M_{\mathcal{C}}(p_h, T')$ .
- (2) Synchronous Euclidean Distance: Under this distance measure,  $p'_h$  can be calculated with the following steps. The first step is to find the segment  $\overline{p_{s_k}p_{s_{k+1}}}$  of T' with  $s_k \leq h \leq s_{k+1}$ . The second step is to find a point along the line passing through two points, namely  $(x_{s_k}, y_{s_k}, t_{s_k})$  and  $(x_{s_{k+1}}, y_{s_{k+1}}, t_{s_{k+1}})$ , in a three-dimensional space such that the third dimensional value (representing the time dimension) of this point is  $t_h$ . Then,  $p'_h$  is set to be the first two-dimensional values of this point. Simi-

larly, we define a mapping function  $M_S$  which maps both  $p_h$  and T' to  $p'_h$ . That is,  $p'_h = M_S(p_h, T')$ .

However, all of them adopt *position-based* distances instead of the *direction-based* distance studied in this thesis. In the next section, we show that they do not give any error guarantee on the direction information.

### **Theoretical Properties**

The *length* between two positions  $p_i$  and  $p_j$  wrt T (i < j), denoted by  $len(p_i, p_j | T)$ , is defined to be the length of the trace from  $p_i$  to  $p_j$  in T. That is,

$$len(p_i, p_j | T) = \sum_{k=i}^{j-1} d(p_k, p_{k+1})$$

The (average) *speed* between two positions  $p_i$  and  $p_j$  wrt T (i < j), denoted by  $speed(p_i, p_j|T)$ , is equal to  $len(p_i, p_j|T)/(t_j - t_i)$ , where  $t_i$  ( $t_j$ ) is the time stamp corresponding to  $p_i$  ( $p_j$ ).

Interestingly, DPTS gives error guarantees on the length and speed information. Consider that T is a trajectory and T' is an  $\epsilon_t$ -simplification of T with  $\epsilon_t < \pi/2$ . For any two adjacent positions  $p_i$  and  $p_{i+1}$  in T where  $i \in [1, n)$ , both the length and the speed between the two corresponding *estimated* positions wrt T' are theoretically bounded. The estimated positions are determined by a mapping function. For the sake of space, in the remaining of the paper, if we do not specify the distance measure, we mean that we adopt the mapping function used in the Closest Euclidean Distance (i.e.,  $M_{\mathcal{C}}(\cdot, \cdot)$ ). The results based on the other mapping function (i.e.,  $M_{\mathcal{S}}(\cdot, \cdot)$ ) can be found in Section A.2 in the appendix.

**Lemma 4.3.1 (Bounded Length/Speed)** Let T be a trajectory and T' be an  $\epsilon_t$ -simplification of T with  $\epsilon_t < \pi/2$ . For any two adjacent positions  $p_i$  and  $p_{i+1}$  in T

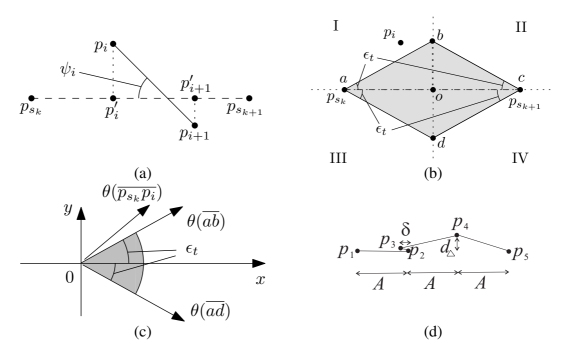


Figure 4.4: Proofs of Lemma 4.3.1, Lemma 4.3.2 and Lemma 4.3.3, DPTS where  $i \in [1, n)$ ,

$$\cos(\epsilon_t) \le \frac{len(p'_i, p'_{i+1}|T')}{len(p_i, p_{i+1}|T)} \le 1 \ and \ \cos(\epsilon_t) \le \frac{speed(p'_i, p'_{i+1}|T')}{speed(p_i, p_{i+1}|T)} \le 1$$

where 
$$p'_i = M_{\mathcal{C}}(p_i, T')$$
 and  $p'_{i+1} = M_{\mathcal{C}}(p_{i+1}, T')$ .

**Proof.** Let  $\overline{p_{s_k}p_{s_{k+1}}}$  be the segment of T' such that  $p_{s_k}$  is the last position with  $s_k \leq i$  and  $p_{s_{k+1}}$  is the first position with  $s_{k+1} \geq i+1$ . Consider Figure 4.4(a) for illustration. Since  $\epsilon_t < \pi/2$ , we can verify that  $p'_i$  and  $p'_{i+1}$  are located along  $\overline{p_{s_k}p_{s_{k+1}}}$ .

Let  $\psi_i$  be the angle formed by the two lines that pass through  $\overline{p_i p_{i+1}}$  and  $\overline{p_{s_k} p_{s_{k+1}}}$ . Thus,

$$len(p'_{i}, p'_{i+1}|T') = d(p'_{i}, p'_{i+1}) = \cos(\psi_{i}) \cdot d(p_{i}, p_{i+1})$$
$$= \cos(\psi_{i}) \cdot len(p_{i}, p_{i+1}|T) \ge \cos(\epsilon_{t}) \cdot len(p_{i}, p_{i+1}|T)$$

which implies that  $\cos(\epsilon_t) \leq \frac{len(p_i',p_{i+1}'|T')}{len(p_i,p_{i+1}|T)} \leq 1$ . Besides, since  $speed(p_i',p_{i+1}'|T') = len(p_i',p_{i+1}'|T')/(t_{i+1}-t_i)$  and  $speed(p_i,p_{i+1}|T) = len(p_i,p_{i+1}|T)/(t_{i+1}-t_i)$ , we know  $\cos(\epsilon_t) \leq \frac{speed(p_i',p_{i+1}'|T')}{speed(p_i,p_{i+1}|T)} \leq 1$ .

Interestingly, DPTS gives an error bound on the position information (in addition to the length/speed information).

**Lemma 4.3.2 (Bounded Position Error)** Let T be a trajectory and T' be an  $\epsilon_t$ simplification of T with  $\epsilon_t < \pi/2$ . For each position  $p_i$  in T where  $i \in [1, n]$ , we have

$$d(p_i, p_i') \le 0.5 \cdot \tan(\epsilon_t) \cdot L_{max}$$

where 
$$p'_{i} = M_{\mathcal{C}}(p_{i}, T')$$
 and  $L_{max} = \max_{k \in [1,m)} len(p_{s_{k}}, p_{s_{k+1}} | T')$ .

**Proof.** Let  $\overline{p_{s_k}p_{s_{k+1}}}$  be the segment of T' such that  $p_{s_k}$  is the last position with  $p_{s_k} \leq i$  and  $p_{s_{k+1}}$  is the first position with  $p_{s_{k+1}} \geq i$ . We construct a rhombus  $\diamond_{abcd}$  with four corners, namely a,b,c and d, such that a is at  $p_{s_k}$ , c is at  $p_{s_{k+1}}$  and the angle between  $\overline{ab}$   $\overline{(cb)}$  and  $\overline{ad}$   $\overline{(cd)}$  is equal to  $2 \cdot \epsilon_t$ . Consider Figure 4.4(b) for illustration where  $\diamond_{abcd}$  is indicated by the shaded area. We claim that  $p_i$  is inside  $\diamond_{abcd}$  which we prove by contradiction.

Assume that  $p_i$  is outside  $\diamond_{abcd}$ . We partition the plane into 4 parts with the two lines that pass through  $\overline{ac}$  and  $\overline{bd}$  as indicated by I, II, III and IV in Figure 4.4(b), where o is the intersection of the two lines. We consider 4 cases of which partition  $p_i$  is in. Without loss of generality, suppose  $p_i$  falls in part I.

Since  $p_i$  is outside  $\diamondsuit_{abcd}$ , we know  $\theta(\overline{p_{s_k}, p_i})$  falls outside range  $[\theta(\overline{ad}), \theta(\overline{ab})]$ . For illustration, consider Figure 4.4(c).

Consider the segments between  $p_{s_k}$  and  $p_i$  in T. For each such segment  $\overline{p_h p_{h+1}}$  ( $s_k \leq h < i$ ), we denote by  $\overline{p_h p_{h+1}}$  the vector from  $p_h$  to  $p_{h+1}$ . We know that the direction of each such vector falls in range  $[\theta(\overline{ad}), \theta(\overline{ab})]$  since otherwise  $\epsilon(T') > \epsilon_t$ . As a result, we know  $\theta(\overline{p_{s_k} p_i})$  falls in range  $[\theta(\overline{ad}), \theta(\overline{ab})]$  since  $\overline{p_{s_k} p_i} = \sum_{s_k \leq h < i} \overline{p_h p_{h+1}}$ . This, however, contradicts the fact that  $\theta(\overline{p_{s_k} p_i})$  falls outside range  $[\theta(\overline{ad}), \theta(\overline{ab})]$ .

Thus, we know that  $p_i$  falls in  $\diamond_{abcd}$ . Therefore, we have

$$d(p_i, p_i') \le d(b, o) = \tan(\epsilon_t) \cdot d(a, o) = 0.5 \cdot \tan(\epsilon_t) \cdot d(a, c)$$
$$= 0.5 \cdot \tan(\epsilon_t) \cdot len(p_{s_k}, p_{s_{k+1}} | T') \le 0.5 \cdot \tan(\epsilon_t) \cdot L_{max}$$

which finishes the proof.

Next, we show that existing position-preserving simplified trajectories do not have bounds on the direction information.

**Lemma 4.3.3 (Unbounded Direction Error)** Let T be a trajectory and T' be a (direction-based)  $\epsilon_t$ -simplification of T with  $\epsilon_t < \pi/2$ . Let  $T_{\mathcal{C}}$  be a (position-based) simplified trajectory of T such that  $|T_{\mathcal{C}}| = |T'|$  and the error of  $T_{\mathcal{C}}$  under the Closest Euclidean Distance is minimized. There exists a dataset such that  $\epsilon(T_{\mathcal{C}}) \approx \pi$  and  $\epsilon(T') \approx 0$ .

**Proof.** We prove by constructing a problem instance as shown in Figure 4.4(d).  $T=(p_1,p_2,p_3,p_4,p_5)$  is a trajectory, where  $p_1,p_2,p_3$  and  $p_5$  are located at a horizontal line and  $p_4$  has its perpendicular distance from this line equal to a small real number  $d_{\triangle}$ . Besides,  $d(p_2,p_3)=\delta$  where  $\delta << d_{\triangle}$ .

Suppose that we can only keep 4 positions in the simplified trajectory. In other words, we have to remove 1 position from the 5 positions. If we consider preserving the direction information,  $p_4$  will be removed and thus  $T'=(p_1,p_2,p_3,p_5)$ . Thus,  $\epsilon(T')=\epsilon(\overline{p_3p_5})\approx 0$ . If we consider preserving the position information,  $p_2$  will be removed and thus  $T_{\mathcal{C}}=(p_1,p_3,p_4,p_5)$ . Hence,  $\epsilon(T_{\mathcal{C}})=\epsilon(\overline{p_1p_3})=\triangle(\theta(\overline{p_1p_3}),\theta(\overline{p_2p_3}))\approx \pi$ .

# 4.3.2 Existing Trajectory Simplification

Many trajectory simplification techniques have been proposed. We categorize them by the main idea employed in the algorithm as follows. They are *Split* [50, 24], *Merge* 

[55, 51], *Greedy* [40, 50] and *Dead-Reckoning* [42]. *Split* is an approach which finds a position in a given trajectory, according to the *heuristic* value of the position, to *split* the whole trajectory into two sub-trajectories and continues the process iteratively on each of the split sub-trajectories which cannot be approximated by a line segment connecting its start position and its end position. *Merge* is an approach which finds two adjacent segments in a given trajectory, according to the *heuristic* value computed from these two adjacent segments, discards the position *p* bridging these two segments, and create a segment connecting the non-bridging end position of one segment and the non-bridging end position of the other segment. It continues the process iteratively until discarding any position *p* violates the error tolerance. *Greedy* is an approach which finds a sequence of the greatest number of consecutive segments to be discarded and create a segment connecting the two end positions of this sequence iteratively until discarding any sequence of 2 consecutive segments violates the tolerance constraint. *Dead-Reckoning* is an online algorithm which reads each position sequentially and determines whether this position is discarded or not according to a *heuristic* criterion.

The aforementioned ideas of *Split*, *Merger* and *Greedy* can be adapted to our DPTS problem. The only change is to change the error measurement to our simplification error (Equation (4.2)). However, they have their drawbacks. First, they cannot return *optimal* solutions. Second, as shown in our experiments, they are not efficient compared with our proposed *Intersect* algorithm. Details of the adaptation can be found in Section A.3 of the appendix.

Other related studies include [9] which studies the error bounds of several queries on the simplified trajectories with bounded simplification errors mainly measured by the position information, [17] which studies the trajectory simplification problem with the consideration of the shape and also the semantic meanings of the trajectory, [16] which introduces a multi-resolution *polygonal curve approximation* (also called *line* 

simplification) algorithm for trajectory simplification, and [39] which studies the trajectory simplification problem where the trajectories are constrained to a road network. None of these studies pay attention to the direction information for trajectory simplification.

## 4.4 Finding Optimal Solution

A naive solution for the DPTS problem is to traverse each possible simplification of T with its simplification error at most  $\epsilon_t$  and then to pick the one with the smallest size. Since the number of all possible simplifications of a trajectory T is  $2^{|T|-2}$ , this solution is not feasible in practice. Alternatively, one may adopt a common dynamic programming (DP) technique for the DPTS problem. Unfortunately, the time complexity of this technique is cubic. For the sake of space, we include this DP algorithm in Section A.4 in the appendix. Instead, we propose a method called SP which is much faster and scalable.

Algorithm SP involves the following three steps.

- **Step 1** (**Graph Construction**): It first constructs a graph based on the given trajectory.
- Step 2 (Shortest Path Finding): It computes a shortest path in this graph.
- **Step 3 (Solution Generation):** It finally returns the solution for DPTS according to the shortest path found.

In Step 1, it constructs a graph wrt  $\epsilon_t$ , denoted by  $G_{\epsilon_t}(V, E)$ , as follows. For each position  $p_i$  of T where  $1 \leq i \leq n$ , it creates a vertex for  $p_i$  in V. For each pair of two positions  $(p_i, p_j)$  where i < j, it creates an edge  $(p_i, p_j)$  in E if  $\epsilon(\overline{p_i p_j}) \leq \epsilon_t$ .

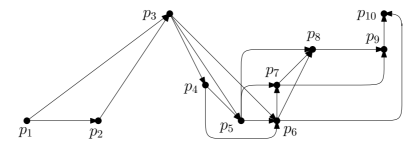


Figure 4.5: The graph  $G_{\epsilon_t}$  constructed based on the running example when  $\epsilon_t$  is set to be  $\pi/4=0.785$ , DPTS

In Step 2, it finds the shortest path from  $p_1$  to  $p_n$  in  $G_{\epsilon_t}$  by a shortest path algorithm (e.g., a BFS search). Here, the *length* of a path is defined to be the number of edges involved along the path.

In Step 3, it generates the solution for DPTS according to the shortest path found. Note that all vertices involved in this shortest path correspond to all positions in the  $\epsilon_t$ -simplification of T with the smallest size. Thus, if the ordering of the positions (or vertices) involved in the shortest path is " $p_{s_1}$ - $p_{s_2}$ -...- $p_{s_m}$ ", it returns the solution T' as  $(p_{s_1}, p_{s_2}, ..., p_{s_m})$ .

**Example 6 (Algorithm SP)** Consider our running example in Figure 4.2. Suppose that  $\epsilon_t = 0.785$ . In Step 1 of the *SP* algorithm, we can construct graph  $G_{\epsilon_t}$  accordingly as shown in Figure 4.5. In this figure, we construct a vertex for each position in T. Besides, for each pair of positions  $p_i$  and  $p_j$  where i < j, if  $\epsilon(\overline{p_i p_j}) \le \epsilon_t$ , we create an edge  $(p_i, p_j)$ . Note that  $\epsilon(\overline{p_i p_{i+1}}) = 0$  for each  $i \in [1, n-1]$ .

In Step 2, we can find the shortest path in this graph. It is easy to verify that  $p_1$ - $p_3$ - $p_6$ - $p_{10}$  is the shortest path. Finally, in Step 3, we construct the solution of DPTS as  $(p_1, p_3, p_6, p_{10})$ .

Let us analyze the time complexity of a *straightforward* implementation of algorithm *SP*. For Step 1, a straightforward solution for constructing  $G_{\epsilon_t}$  is to try all possible pairs of  $(p_i, p_j)$  where  $1 \leq i < j \leq n$  and to check whether  $\epsilon(\overline{p_i p_j}) \leq \epsilon_t$ .

### **Algorithm 7** The SP algorithm with the practical enhancement

```
Require: A trajectory T = (p_1, p_2, ..., p_n) and the error tolerance \epsilon_t
  1: H_0 \leftarrow \{p_1\}; U \leftarrow \{p_2, p_3, ..., p_n\}; l \leftarrow 1
  2:
3:
      while true do H_l \leftarrow \emptyset
          //process the positions in H_{l-1} and U in a reversed order for each p_i in H_{l-1} and each p_j in U where i < j do
  4:
  5:
              if \epsilon(\overline{p_ip_i}) \leq \epsilon_t then
  6:
  7:
                  if p_i = p_n then
                      return the trajectory corresponding to the shortest path from p_1 to p_n
  8:
                   U \leftarrow U \setminus \{p_i\}; H_l \leftarrow H_l \cup \{p_i\}
  9:
          l \leftarrow l + 1
10:
```

Since there are  $O(n^2)$  possible such pairs and the checking cost for each pair is O(n), the time complexity of Step 1 is  $O(n^3)$ . For Step 2, a simple BFS could be adopted to find the shortest path from  $p_1$  to  $p_n$  in  $G_{\epsilon_t}$ , which takes O(|V| + |E|) time. Since |V| = O(n) and  $|E| = O(n^2)$ , we know that the cost of BFS is  $O(n^2)$ . Step 3 which returns the solution takes O(n) time. As we can see, the time complexity of Step 1 (i.e., the graph construction) dominates those of Step 2 and Step 3. Thus, the overall time complexity of a straightforward implementation of algorithm SP is  $O(n^3)$ . Besides, the space complexity of SP is simply O(|V| + |E|) which corresponds to the space cost of maintaining  $G_{\epsilon_t}$ .

In the following, we propose two kinds of enhancement techniques in order to improve the efficiency of our *SP* algorithm. The first one is called the *practical enhancement* (Section 4.4.1) which is to improve the performance of the algorithm in a practical way. The second one is called the *complexity improvement* (Section 4.4.2) which is to improve the theoretical time complexity of the algorithm from cubic to quadratic with some properties.

### 4.4.1 Practical Enhancement

The practical enhancement is to construct  $G_{\epsilon_t}$  (in Step 1) on the demand of the BFS procedure used in Step 2. Specifically, the straightforward implementation has to materialize all possible edges in  $G_{\epsilon_t}$  in Step 1 and then perform the BFS procedure

in Step 2. Here, the enhancement constructs only *some* of the edges in  $G_{\epsilon_t}$  which are needed in the BFS procedure. Since some other edges need not be constructed, the space consumption can be reduced and some computations can be also saved.

Given a position p in T and a non-negative integer l, p is said to be an l-length position if the length of the shortest path from  $p_1$  to p in  $G_{\epsilon_t}$  is equal to l. Given a non-negative integer l, we define the l-length unique set, denoted by  $H_l$ , to be the set of all l-length positions in T. For example, the 0-length unique set  $H_0$  is  $\{p_1\}$ . Consider the BFS procedure starting from  $p_1$  on  $G_{\epsilon_t}$ . It first retrieves the set of positions which are the out-neighbors of  $p_1$ . This set corresponds to  $H_1$ . Then, starting from each position p in  $H_1$ , it retrieves the set of positions which are the out-neighbors of p and have not been retrieved before. This set corresponds to  $H_2$ . The above process continues from  $H_2$  in the same manner until  $p_n$  is retrieved.

In view of the above discussion, we design our SP algorithm with this enhancement as follows. We maintain the l-length unique sets  $H_l$  (l=0,1,2,...) which store the positions retrieved by the BFS procedure and U for storing the remaining positions that have not been retrieved by the BFS procedure. We initialize  $H_0$  to be  $\{p_1\}$  and U to be  $\{p_2,p_3,...,p_n\}$ . We then compute  $H_l$  based on  $H_{l-1}$  for l=1,2,... iteratively as follows. We start from each position  $p_i$  in  $H_{l-1}$ . For each position  $p_j$  in U, we compute  $\epsilon(\overline{p_ip_j})$ . If  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$ , we further check whether  $p_j$  is  $p_n$ . If so, we stop the process since the shortest path from  $p_1$  to  $p_n$  has been found; otherwise, we exclude  $p_j$  from U and include it in  $H_l$ . Besides, when processing the positions in  $H_{l-1}$  and U, we impose a reversed order, which corresponds to  $p_n, p_{n-1}, ..., p_1$ . The intuition is that we expect that  $p_n$  could be retrieved earlier in this way. We present our enhanced SP algorithm in Algorithm 7.

**Complexity Analysis.** The worst-case time complexity of the *SP* algorithm with the practical enhancement keeps the same as that of the straightforward implementation,

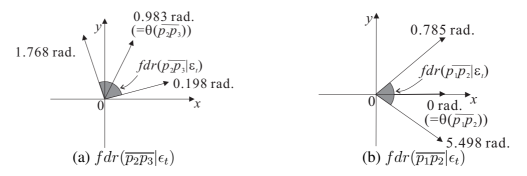


Figure 4.6: Illustration of  $fdr(\overline{p_2p_3}|\epsilon_t)$  and  $fdr(\overline{p_1p_2}|\epsilon_t)$ , DPTS

i.e., it is still  $O(n^3)$ . However, in practice, with the practical enhancement, the SP algorithm is more efficient since some computations of  $\epsilon(\overline{p_ip_j})$  are avoided, and it is also more scalable since there is no need to materialize  $G_{\epsilon_t}$ . The space complexity of SP with the practical enhancement is simply O(n) since it maintains each position once and does not materialize  $G_{\epsilon_t}$  explicitly.

## 4.4.2 Complexity Improvement

The complexity improvement is to improve the time complexity of our *SP* algorithm from cubic to quadratic by using some properties for our algorithm. In this section, we focus on the complexity improvement based on the straightforward implementation for illustration. In Section 4.4.3, we describe how this complexity improvement can be incorporated with the practical enhancement.

As can be noticed, the cost of the straightforward implementation is dominated by the construction of graph  $G_{\epsilon_t}$ . In this section, we propose a technique to reduce the cost of constructing the graph from  $O(n^3)$  to  $O(C \cdot n^2)$  time, where C is shown to be a small constant in most cases. The major idea of such an improvement is to reduce the time complexity of checking whether  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$  (in the graph construction step) from O(n) to O(C) by utilizing a new concept called "feasible direction range". Before we present the main idea, we first introduce some related concepts.

Given two angles  $\theta_1$  and  $\theta_2$  in  $[0, 2\pi)$ , an angular range, represented in the form

of  $[\theta_1, \theta_2]$ , is defined to be a set of all possible angles of a vector originated from the origin when it is rotated anti-clockwise from  $\theta_1$  to  $\theta_2$ . For example, the shaded part in Figure 4.6(a) shows the angular range of [0.198, 1.768], and the shaded part in Figure 4.6(b) shows the angular range of [5.498, 0.785]. Since the direction of  $\overline{p_2p_3}$  is 0.983 radian, we say that its direction is in [0.198, 1.768] but not in [5.498, 0.785]. Similarly, since the direction of  $\overline{p_1p_2}$  is 0 radian, we say that its direction is in [5.498, 0.785] but not in [0.198, 1.768].

**Definition 4.4.1 (feasible direction range)** Given a segment  $\overline{p_h p_{h+1}}$   $(1 \le h < n)$  in T, the **feasible direction range** of  $\overline{p_h p_{h+1}}$  wrt  $\epsilon_t$ , denoted by  $fdr(\overline{p_h p_{h+1}} | \epsilon_t)$ , is defined to be the angular range in the form of  $[\theta_1, \theta_2]$  with  $\theta_1 = [(\theta(\overline{p_h p_{h+1}}) - \epsilon_t) \mod 2\pi]$  and  $\theta_2 = [(\theta(\overline{p_h p_{h+1}}) + \epsilon_t) \mod 2\pi]$ .

The feasible direction range of  $\overline{p_h p_{h+1}}$  wrt  $\epsilon_t$  corresponds to a set of all possible directions each of which has its angular difference from  $\overline{p_h p_{h+1}}$  at most  $\epsilon_t$ . We can write fdr as follows.

$$fdr(\overline{p_h p_{h+1}}|\epsilon_t) = [\theta(\overline{p_h p_{h+1}}) - \epsilon_t, \theta(\overline{p_h p_{h+1}}) + \epsilon_t] \mod 2\pi$$
 (4.3)

Consider our running example. Suppose that  $\epsilon_t$  is set to 0.785. Since the direction of  $\overline{p_2p_3}$  is 0.983 radian,  $fdr(\overline{p_2p_3}|\epsilon_t)=([0.983-0.785,0.983+0.785]\mod 2\pi)=[0.198,1.768]$  (See Figure 4.6(a)). Similarly, since the direction of  $\overline{p_1p_2}$  is 0 radian,  $fdr(\overline{p_1p_2}|\epsilon_t)=([0-0.785,0+0.785]\mod 2\pi)=[5.498,0.785]$  (See Figure 4.6(b)).

We denote by T[i,j] the sub-trajectory of T that is between position  $p_i$  and position  $p_j$  ( $1 \le i < j \le n$ ), i.e.,  $T[i,j] = (p_i, p_{i+1}, ..., p_j)$ . We define the *feasible direction* range of a sub-trajectory T[i,j] wrt  $\epsilon_t$ , denoted by  $fdr(T[i,j]|\epsilon_t)$ , to be the *intersection* of the fdr's of the segments in T[i,j]. That is,

$$fdr(T[i,j]|\epsilon_t) = \bigcap_{i \le h < j} fdr(\overline{p_h p_{h+1}}|\epsilon_t)$$
(4.4)

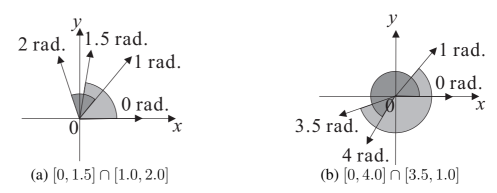


Figure 4.7: Illustration of intersection operations between two angular ranges, DPTS

In our running example,  $fdr(T[1,3]|\epsilon_r)$  is equal to  $\bigcap_{1 \leq h < 3} fdr(\overline{p_h p_{h+1}}|\epsilon_t) = fdr(\overline{p_1 p_2}|\epsilon_t) \cap fdr(\overline{p_2 p_3}|\epsilon_t) = [5.498, 0.785] \cap [0.198, 1.768] = [0.198, 0.785]$ . Figures 4.6(a) and (b) illustrate this scenario.

In the following, we simply write  $fdr(\overline{p_hp_{h+1}})$  (fdr(T[i,j])) for  $fdr(\overline{p_hp_{h+1}}|\epsilon_t)$   $(fdr(T[i,j]|\epsilon_t))$  if the context of  $\epsilon_t$  is clear.

In some cases, an intersection between an angular range and another angular range results in a *single* angular range. In some other cases, this intersection operation results in *multiple* disjoint angular ranges. To illustrate, an intersection between [0, 1.5] and [1.0, 2.0] results in a single angular range [1, 1.5] (as shown in Figure 4.7(a)). An intersection between [0, 4.0] and [3.5, 1.0] results in two angular ranges [3.5, 4.0] and [0, 1.0] (as shown in Figure 4.7(b)).

Thus, from Equation (4.4), since fdr(T[i,j]) involves multiple intersection operations of angular ranges, it may consist of multiple disjoint angular ranges. We denote by ||fdr(T[i,j])|| the number of disjoint angular ranges in fdr(T[i,j]).

With the concept of "feasible direction range", we are now ready to describe how we can check whether  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$  efficiently.

**Lemma 4.4.1** Let 
$$T=(p_1,p_2,...,p_n)$$
 be a trajectory.  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$  iff  $\theta(\overline{p_ip_j})$  is in  $fdr(T[i,j])$ .

**Proof.** "if": Assume that  $\theta(\overline{p_ip_j})$  is in fdr(T[i,j]). It follows that  $\theta(\overline{p_ip_j})$  is in

 $fdr(\overline{p_hp_{h+1}})$  and thus  $\triangle(\theta(\overline{p_ip_j}), \theta(\overline{p_hp_{h+1}})) \le \epsilon_t$  for  $i \le h < j$ . Therefore,  $\epsilon(\overline{p_ip_j}) = \max\{\triangle(\theta(\overline{p_hp_{h+1}}), \theta(\overline{p_ip_j}))|i \le h < j\} \le \epsilon_t$ .

"only-if": this direction could be verified similarly. ■

Lemma 4.4.1 suggests that checking whether  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$  is equivalent to checking whether  $\theta(\overline{p_ip_j})$  is in fdr(T[i,j]). Suppose that fdr(T[i,j]) has been computed. Then, checking whether  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$  takes O(||fdr(T[i,j])||) only (compared with O(n) in the straightforward implementation).

Note that in some cases,  $\epsilon(\overline{p_ip_j}) > \epsilon_t$  but  $\epsilon(\overline{p_ip_{j+k}}) \le \epsilon_t$  where j > i > 0 and k > 0. By Lemma 4.4.1, we know that  $\theta(\overline{p_ip_j})$  is not in fdr(T[i,j]) but  $\theta(\overline{p_ip_{j+k}})$  is in fdr(T[i,j+k]). To illustrate, in our running example (Figure 4.2), if we set  $\epsilon_t = \pi/4$ , then  $\epsilon(\overline{p_6p_9}) > \epsilon_t$  but  $\epsilon(\overline{p_6p_{10}}) \le \epsilon_t$ . By Lemma 4.4.1,  $\theta(\overline{p_6p_9})$  is not in fdr(T[6,9]) but  $\theta(\overline{p_6p_{10}})$  is in fdr(T[6,10]).

Now, we know that the checking step can be done in O(||fdr(T[i,j])||). There are two remaining issues related to this checking step. The first issue is related to the size of fdr(T[i,j]). If this size is very large, the checking step is still expensive. Fortunately, we find that this size is usually a small constant. When  $\epsilon_t \leq \pi/2$ , it is equal to 1. The second issue is how to compute fdr(T[i,j]) efficiently for each i and j where i < j.

**Issue 1: Size of** fdr(T[i, j])

**Lemma 4.4.2** Let  $T=(p_1,p_2,...,p_n)$  be a trajectory and  $\epsilon_t$  be the error tolerance. Then, given two integers i and j  $(1 \leq i < j \leq n)$ , ||fdr(T[i,j])|| is bounded by  $\min\{1+\lfloor \frac{\epsilon_t}{(\pi-\epsilon_t)}\rfloor,j-i\}$ .

**Proof.** We first give some concepts based on an angular range and provide a lemma which is used to prove Lemma 4.4.2.

We denote the *universe* angular range  $[0, 2\pi)$  by  $\mathcal{U}$ , Given an angular range [a, b], we denote its *complement* wrt  $\mathcal{U}$ , which is the angular range (b, a), by  $[a, b]^c$ . We define the *span* of an angular range [a, b], denoted by [a, b].span, to be equal to  $(b - a)(\mod 2\pi)$ . We define the *span* of fdr(T[i, j])  $(1 \le i < j \le n)$ , denoted by fdr(T[i, j]).span, to be equal to the sum of the spans of the disjoint angular ranges that are involved in fdr(T[i, j]).

**Lemma 4.4.3** Let [a,b] and [a',b'] be two angular ranges.  $[a,b] \cap [a',b']$  involves two disjoint angular ranges iff  $[a',b']^c$  (i.e., (b',a')) falls in [a,b] completely.

**Proof.** This could be verified easily by the fact that  $[a,b] \cap [a',b']$  involves two disjoint angular ranges, namely [a,b'] and [a',b], iff (b',a') falls in [a,b] completely.

Suppose  $fdr(\overline{p_kp_{k+1}})$  is  $[a_k,b_k]$  for  $i\leq k< j$ . Then, fdr(T[i,j]) is equal to  $\bigcap_{k=i}^{j-1}[a_k,b_k]$ . Besides, we know  $[a_k,b_k].span=2\epsilon_t$  for  $i\leq k< j$  since  $a_k=\theta(\overline{p_kp_{k+1}})-\epsilon_t \mod 2\pi$  and  $b_k=\theta(\overline{p_kp_{k+1}})+\epsilon_t \mod 2\pi$  (Definition 4.4.1).

First, we prove  $||fdr(T[i,j])|| \leq j-i$  by induction of k=j-i. Base step: k=1. The correctness is obvious since  $fdr(T[i,i+1])=[a_i,b_i]$  which involves one angular range only (i.e.,  $[a_i,b_i]$ ) and thus ||fdr(T[i,i+1])||=1. Induction step:  $||fdr(T[i,i+k)|| \leq k$  implies  $||fdr(T[i,i+k+1)|| \leq k+1$ . Assume ||fdr(T[i,i+k)|| = r ( $r \leq k$ ). Specifically, let  $[a'_1,b'_1], [a'_2,b'_2], ..., [a'_r,b'_r]$  be the r disjoint angular ranges involved in fdr(T[i,i+k]). Then, fdr(T[i,i+k+1]), which is equal to  $fdr(T[i,i+k]) \cap [a_{i+k},b_{i+k}]$ , corresponds to r intersections,  $[a'_h,b'_h] \cap [a_{i+k},b_{i+k}]$  for  $1 \leq h \leq r$ , each two of which are disjoint. Among these r intersections, we show that at most one involves two disjoint angular ranges by contradiction. Assume that there exists  $h_1$  and  $h_2$  ( $i \leq h_1 \neq h_2 \leq r$ ) such that both  $[a'_{h_1},b'_{h_1}] \cap [a_{i+k},b_{i+k}]$  and  $[a'_{h_2},b'_{h_2}] \cap [a_{i+k},b_{i+k}]$  involve two disjoint angular ranges. According to Lemma 4.4.3, we know that  $(b_{i+k},a_{i+k})$  is in both  $[a'_{h_1},b'_{h_1}]$  and  $[a'_{h_2},b'_{h_2}]$ , which leads to a contradiction since

 $[a'_{h_1},b'_{h_1}]$  and  $[a'_{h_2},b'_{h_2}]$  are disjoint. As a result,  $fdr(T[i,i+k_1])$  involves at most r+1 disjoint angular ranges. That is,  $||fdr(T[i,i+k+1])|| \le r+1 \le k+1$ .

Second, we prove  $||fdr(T[i,j])|| \leq 1 + \lfloor \frac{\epsilon_t}{\pi - \epsilon_t} \rfloor$ . We compute fdr(T[i,i+1]), fdr(T[i,i+2]), ..., fdr(T[i,j]) sequentially based on the following equation.

$$fdr(T[i, i+k+1]) = fdr(T[i, i+k]) \cap [a_{i+k}, b_{i+k}]$$

$$= fdr(T[i, i+k]) \setminus (b_{i+k}, a_{i+k})$$
(4.5)

We have two cases regarding Equation 4.5. Case 1: ||fdr(T[i,i+k+1])|| = ||fdr(T[i,i+k])|| + 1. In this case, according to Lemma 4.4.3,  $(b_{i+k},a_{i+k})$  is in one of the disjoint angular ranges that are involved in fdr(T[i,i+k+1]). Then, we deduce that  $fdr(T[i,i+k+1]).span = fdr(T[i,i+k]).span - (2\pi - 2\epsilon_t)$  since  $(a_{i+k} - b_{i+k})$  mod  $2\pi = 2\pi - 2\epsilon_t$ . Case 2: ||fdr(T[i,i+k+1])|| = ||fdr(T[i,i+k])||. In this case, we have  $fdr(T[i,i+k+1]).span \le fdr(T[i,i+k]).span$  since the span is non-increasing after an intersection operation. In view of the above two cases, we conclude that the *increase* of the number of disjoint angular ranges by 1 (in Case 1 only) is due to the *decrease* of the span by  $(2\pi - 2\epsilon_t)$ . Since at the beginning, ||fdr(T[i,i+1])|| = 1 and  $fdr(T[i,i+1]).span = 2\epsilon_t$ , ||fdr(T[i,i+k])|| has its greatest value equal to  $(1+\lfloor \frac{2\epsilon_t}{2\pi - 2\epsilon_t}\rfloor)$ .

According to Lemma 4.4.2, ||fdr(T[i,j])|| is usually bounded by a small constant. Let  $C = \min\{1 + \lfloor \frac{\epsilon_t}{(\pi - \epsilon_t)} \rfloor, j - i\}$ . In particular, when  $\epsilon_t \leq \pi/2$ ,  $\lfloor \frac{\epsilon_t}{(\pi - \epsilon_t)} \rfloor$  is equal to 0. In this case, ||fdr(T[i,j])|| is exactly equal to  $\min\{1, j - i\} = 1$  (since j > i).

### **Issue 2: How to Compute** fdr(T[i, j]) **Efficiently**

A straightforward method to compute fdr(T[i,j]) is to compute  $fdr(\overline{p_hp_{h+1}})$  ( $i \leq h < j$ ) independently and then to intersect these fdr's. This straightforward method, nevertheless, incurs the cost of  $\Omega(n)$  on average. Since we have  $\Theta(n^2)$  instances of

fdr(T[i,j]), computing all instances of fdr(T[i,j]) with this method incurs the total cost of  $\Omega(n^3)$ . Fortunately, this method could be improved significantly since it involves a lot of redundant work. A better method only takes O(C) instead of  $\Omega(n)$  to compute fdr(T[i,j]) based on the following incremental property.

Given two integers i and j where  $1 \le i < j < n$ ,

$$fdr(T[i,j+1]) = fdr(T[i,j]) \cap fdr(\overline{p_i p_{i+1}})$$
(4.6)

Suppose that the content of fdr(T[i,j]) is known, since the total number of angular ranges in  $fdr(\overline{p_jp_{j+1}})$  is 1, we can compute fdr(T[i,j+1]) in O(C) time. Note that C is the greatest number of angular ranges in fdr(T[i,j]) and the intersection operation between two intervals could be finished in O(1) time.

Thus, we propose to compute fdr(T[i,j])  $(1 \le i < j \le n)$  using the incremental property. Specifically, it involves n rounds.

- At round 1, it computes fdr(T[h, h+1]) (i.e.,  $fdr(\overline{p_hp_{h+1}})$ ) for  $1 \le h < n$ .
- At round r (r > 1), it computes fdr(T[h, h+r]) for  $1 \le h \le n-r$ . Specifically, it computes fdr(T[h, h+r]) by intersecting fdr(T[h, h+r-1]) (which has been maintained at round r-1) and  $fdr(\overline{p_{h+r-1}p_{h+r}})$ . Note that this operation takes O(C) time.

Complexity Analysis. The time complexity of the above method for computing fdr(T[i,j])'s for  $1 \le i < j \le n$  is  $O(C \cdot n^2)$  since it involves n rounds and each round incurs the cost of  $O(C \cdot n)$ . Since there are  $O(n^2)$  times of checking whether  $\epsilon(\overline{p_ip_{i+1}}) \le \epsilon_t$  and each checking can be done in O(C) time with the fdr(T[i,j]) information, the time complexity of the SP algorithm with the complexity improvement is  $O(C \cdot n^2)$ . Besides, the above method of computing fdr(T[i,j])'s has its space complexity of  $O(C \cdot n)$  since at each round, it is sufficient to maintain O(n)

fdf(T[i,j])'s each of which involves O(C) intervals. Since the SP algorithm with the complexity improvement materializes  $G_{\epsilon_t}$  explicitly, we know that its space complexity is  $O(C \cdot n + |V| + |E|)$ , where V(E) is the vertex (edge) set of  $G_{\epsilon_t}$ .

## 4.4.3 Combining The Two Enhancements

The practical enhancement involves the construction of only *some* edges, which means that we just need to perform the checking of  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$  for *some* pairs of positions  $(p_i,p_j)$  only where  $1 \leq i < j \leq n$ . However, the cost of checking whether  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$  is O(n) which is expensive. In contrast, the complexity improvement reduces the cost of checking whether  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$  from O(n) to O(C), but the undesired part is that it always computes fdr(T[i,j]) for *all* pairs of positions  $(p_i,p_j)$ .

In this part, we propose to unify the good aspects of both the practical enhancement and the complexity improvement and at the same time, to avoid their undesired aspects. Our strategy is to maintain fdr(T[i,j])  $(1 \le i < j \le n)$  (i.e., the idea of complexity improvement) on the demand of the BFS process (i.e., the idea of the practical enhancement). Specifically, when checking whether  $\epsilon(\overline{p_ip_j}) \le \epsilon_t$   $(1 \le i < j \le n)$ , we have two cases. Case 1: fdr(T[i,j]) has been computed. In this case, the checking could be finished in O(C) time. Case 2: fdr(T[i,j]) has not been computed. In this case, we recursively resort to fdr(T[i,j-1]) for computing fdr(T[i,j]). This recursive process stops either in Case 1 or fdr(T[i,i+1]) is acquired. Note that fdr(T[i,i+1]) could be computed in O(1) time by Equation (4.3).

This version of SP algorithm enjoys the benefit of the practical enhancement since it checks whether  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$  ( $1 \leq i < j \leq n$ ) on demand of the BFS process and does not materialize  $G_{\epsilon_t}$  explicitly, and it also enjoys the benefit of the complexity improvement since it adopts the "feasible direction range" concept for checking whether  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$ , which is fast.

It could be verified that the worse-case time complexity of the SP algorithm with both the practical enhancement and the complexity improvement is  $O(C \cdot n^2)$  since it computes only a sub-set of all possible fdr(T[i,j])'s.

Besides, the space complexity of this version of SP is  $O(C \cdot n)$  (it is sufficient to maintain for each i the computed fdr[i,j] with the  $largest\ j$  among all computed fdr[i,j]'s throughout the execution of the algorithm since each fdr[i,j] is enquired at most once and note that it does not materialize  $G_{\epsilon_t}$ ).

# 4.5 Finding Approximate Solution

According to the discussion in Section 4.4, the time complexity of an optimal algorithm for DPTS is at least quadratic. This, however, is not scalable enough when the datasets involves millions of positions. In this section, we develop an approximate algorithm called *Intersect* for the DPTS problem, which runs in linear time and gives a certain degree of quality guarantee.

Before we describe *Intersect*, we first give a concept of "feasibility" used in the algorithm. Given an error tolerance  $\epsilon_t$  and two integers i and j where  $1 \leq i < j \leq n$ ,  $\overline{p_i p_j}$  is said to be  $\epsilon_t$ -feasible iff  $fdr(T[i,j]|\epsilon_t)$  is non-empty. With this concept, we have the following property.

**Lemma 4.5.1 (Feasibility)** Given an error tolerance  $\epsilon_t$  and two integers i and j where  $1 \le i < j \le n$ , if  $\overline{p_i p_j}$  is  $\frac{\epsilon_t}{2}$ -feasible, then  $\epsilon(\overline{p_i p_j}) \le \epsilon_t$ .

**Sketch.** Since  $\overline{p_ip_j}$  is  $\frac{\epsilon_t}{2}$ -feasible,  $fdr(T[i,j]|\epsilon_t/2)$  is non-empty. We deduce that for any two segments between  $p_i$  and  $p_j$  in T,  $\overline{p_hp_{h+1}}$  and  $\overline{p_{h'}p_{h'+1}}$  where  $i \leq h < h' < j$ ,  $\triangle(\theta(\overline{p_hp_{h+1}}), \theta(\overline{p_{h'}p_{h'+1}})) \leq \epsilon_t$ . We further deduce that  $\theta(\overline{p_ip_j})$  is in  $fdr(T[i,j]|\epsilon_t)$  because  $\theta(\overline{p_ip_j})$  is equal to the angle of a *vector* which is equal to the sum of all *vectors* between  $p_i$  and  $p_j$  in T (i.e.,  $\sum_{k=i}^{j-1} \overline{p_kp_{k+1}}$  where  $\overline{p_kp_{k+1}}$  is a vector from  $p_k$  to  $p_{k+1}$ 

### Algorithm 8 The Intersect Algorithm

```
Require: A trajectory T=(p_1,p_2,...,p_n); an error tolerance \epsilon_t

Ensure: An \epsilon_t-simplification of T'

1: T' \leftarrow (p_1); e \leftarrow 1; h \leftarrow 2

2: while h \leq n do

3: while h \leq n and \overline{p_e p_h} is \frac{\epsilon_t}{2}-feasible do

4: increment h by 1

5: append p_{h-1} to T'; e \leftarrow h-1

6: return T'
```

for each  $k \in [1, j-1]$ ) in the two-dimensional space (i.e., the *x*-coordinate and the *y*-coordinate). Here, the angle of a vector is defined to be the angle of an anticlockwise rotation from the positive x-axis to this vector. By Lemma 4.4.1,  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$ . The complete proof could be found in Section A.5 in the appendix.

Specifically, *Intersect* has the following steps. Let T' be a variable storing the simplified trajectory to be returned. Let e be a variable storing the position index of the last position in T'. Let h be a variable storing the position index of the position in T being processed. Initially, *Intersect* initializes T' to be  $(p_1)$ , and then sets e to be 1 (since  $p_1$  is currently the last position in T'). Then, it reads each of the remaining positions sequentially. It sets h to 2 (since  $p_2$  is the position in T to process next). It proceeds with an iterative step as follows. Whenever  $h \leq n$  and  $\overline{p_ep_h}$  is  $\frac{\epsilon_t}{2}$ -feasible, it increments h by 1. It terminates this iterative step when either (1) h > n or (2) h has  $f(t) = \frac{\epsilon_t}{2}$ -feasible incremented to a value such that  $f(t) = \frac{\epsilon_t}{2}$ -feasible. For both stopping conditions, we know that  $f(t) = \frac{\epsilon_t}{2}$ -feasible and thus by Lemma 4.5.1, we have  $f(t) = \frac{\epsilon_t}{2}$ . Thus,  $f(t) = \frac{\epsilon_t}{2}$  has above iterative step whenever  $f(t) = \frac{\epsilon_t}{2}$ . Then,  $f(t) = \frac{\epsilon_t}{2}$  has a variable storing the position in  $f(t) = \frac{\epsilon_t}{2}$ . Then,  $f(t) = \frac{\epsilon_t}{2}$  has a variable storing the position in  $f(t) = \frac{\epsilon_t}{2}$ . Then,  $f(t) = \frac{\epsilon_t}{2}$  has a variable storing the position in  $f(t) = \frac{\epsilon_t}{2}$ . Then,  $f(t) = \frac{\epsilon_t}{2}$  has a variable storing the position in  $f(t) = \frac{\epsilon_t}{2}$ . Then,  $f(t) = \frac{\epsilon_t}{2}$  has a variable storing the position in  $f(t) = \frac{\epsilon_t}{2}$ . Then,  $f(t) = \frac{\epsilon_t}{2}$  has a variable storing the position in  $f(t) = \frac{\epsilon_t}{2}$ . Then,  $f(t) = \frac{\epsilon_t}{2}$  has a variable storing the position in  $f(t) = \frac{\epsilon_t}{2}$ . The pseudo-code of  $f(t) = \frac{\epsilon_t}{2}$  has a variable storing the position in  $f(t) = \frac{\epsilon_t}{2}$ . The pseudo-code of  $f(t) = \frac{\epsilon_t}{2}$  has a variable storing the position in  $f(t) = \frac{\epsilon_t}{2}$ . The pseudo-code of  $f(t) = \frac{\epsilon_t}{2}$  has a variable storing the position in  $f(t) = \frac{\epsilon_t}{2}$  has a variable storing the position in  $f(t) = \frac{\epsilon_t}{2}$  has a variable storing the position in  $f(t) = \frac{\epsilon_t}$ 

With Lemma 4.5.1, it is easy to verify that the trajectory returned by *Intersect* is an  $\epsilon_t$ -simplification of T.

**Lemma 4.5.2** Let T' be the output of the Intersect algorithm in Algorithm 8. Then, T' is an  $\epsilon_t$ -simplification of T.

Intersect not only scans the data once only and returns an  $\epsilon_t$ -simplification of T at the end, but also provides a certain degree of guarantee on the size of the simplified trajectory.

**Lemma 4.5.3 (Size Bound)** Let T' be the output of the Intersect algorithm in Algorithm 8. We have  $|T'| \leq |T''|$ , where T'' is the  $\epsilon_t/2$ -simplification of T with the minimum size.

**Sketch.** Let  $T=(p_1,...,p_n)$ ,  $T'=(p_{s_1},...,p_{s_m})$  and  $T''=(p_{r_1},...,p_{r_l})$ . By definition, we have  $s_1=r_1=1$  and  $s_m=r_l=n$ . Note that n=|T|,m=|T'| and l=|T''|. Our proof has two steps. First, we prove that  $fdr(T[r_k,r_{k+1}]|\epsilon_t/2)$  is non-empty for  $1\leq k< l$ . This could be verified by the fact that T'' is an  $\epsilon_t/2$ -simplification. Second, we prove  $m\leq l$  by contradiction which main idea is that if m>l, we show  $s_k\geq r_k$  for k=1,2,...,l by induction, which, however, results in that  $s_m>n$  contradicting  $s_m=n$ . The complete proof could be found in Section A.6 in the appendix.

Complexity Analysis. We know that variable h is incremented whenever line 4 of Algorithm 8 is executed. Both the stopping condition of the outer while-loop (line 2) and one of the stopping conditions of the inner while-loop (line 3) are " $h \le n$ ". We conclude that there are O(n) times to execute the steps in line 3, line 4 and line 5. The step in line 4 and the step in line 5 take O(1) time. In the following, we show that the step in line 3 also takes O(1) time. Thus, the time complexity of *Intersect* is O(n).

The remaining issue is to derive the time complexity of the step in line 3. In line 3, checking whether  $h \leq n$  can be done in O(1) time. However, a straightforward implementation of checking whether  $\overline{p_e p_h}$  is  $\frac{\epsilon_t}{2}$ -feasible (line 3) (or equivalently checking whether  $fdr(T[e,h]|\epsilon_t/2)$  is non-empty) is expensive. This is because as described in

Section 4.4.2, computing  $fdr(T[e,h]|\epsilon_t/2)$  from scratch is expensive. By using the same technique in Section 4.4.2, we can perform this checking operation in O(1) time. Specifically, we introduce a variable called fdr to store the content of  $fdr(T[e,h]|\epsilon_t/2)$ . We have the following two changes in the algorithm due to this variable. Firstly, between line 2 and line 3, we insert a statement that "fdr  $\leftarrow fdr(\overline{p_ep_h}|\epsilon_t/2)$ ". Note that h=e+1 at this moment, and the time complexity of this statement is O(C') where  $C'=\min\{1+\lfloor\frac{\epsilon_t/2}{(\pi-\epsilon_t/2)}\rfloor,h-e\}$  (by using Lemma 4.4.2). Secondly, in the inner while-loop, just after line 4, we insert a statement that "fdr  $\leftarrow$  fdr  $\cap fdr(\overline{p_{h-1}p_h}|\epsilon_t/2)$ ". Note that the time complexity of this statement is O(C'). With these two changes, checking whether  $\overline{p_ep_h}$  is  $\frac{\epsilon_t}{2}$ -feasible (line 3) is equivalent to checking whether the current content of fdr is non-empty because the current content of fdr is equal to  $fdr(T[e,h]|\epsilon_t/2)$  (by Equation (4.6)). Furthermore, since  $\epsilon_t$  is at most  $\pi$  and thus  $\epsilon_t/2$  is at most  $\pi/2$ , we deduce that C'=O(1). We conclude that the time complexity of the step in line 3 is O(1).

The space complexity of *Intersect* is O(n) (which corresponds to the memory usage for storing the simplified trajectory T').

# 4.6 Empirical Studies

## 4.6.1 Datasets and Algorithms

We used 5 real datasets in our experiments, namely Deer, Elk, Hurricane, Geolife and T-Drive. Deer and Elk<sup>1</sup> are two animal movement datasets which contain the radio-telemetry locations of deers in 1995 and elks in 1993, respectively. Hurricane<sup>2</sup> contains the trajectories of the Atlantic hurricanes from year 1950 to year 2004. These three datases (i.e., Deer, Elk and Hurricane) are benchmark datasets for trajectory cluster-

<sup>&</sup>lt;sup>1</sup>http://www.fs.fed.us/pnw/starkey/data/tables/

<sup>&</sup>lt;sup>2</sup>http://weather.unisys.com/hurricane/atlantic/

	# of trajectories	total # of positons	average # of positions per trajectory	directional difference between two adjacent segments (Mean, S.D.)
Deer	32	20,065	627	(1.669, 0.948)
Elk	33	47,204	1,430	(1.647, 0.984)
Hurricane	570	17,736	31	(0.213, 0.300)
Geolife	17,621	24,876,978	1,412	(0.364, 0.615)
T-Drive	10,359	17,740,902	1,713	(0.657, 0.803)

Table 4.1: Real datasets (for DPTS)

ing [46]. Geolife<sup>3</sup> records the outdoor movements of 182 users in a period of 5 years and T-Drive<sup>4</sup> is a set of taxi trajectories in Beijing. These two datasets are widely-used for a broad range of applications on trajectory data [74, 70]. The statistics of these datasets are summarized in Table 4.1.

We study 5 optimal algorithms with the following notions. DP is the dynamic programming algorithm and SP is the straightforward implementation of the *SP* algorithm. SP-prac (SP-theo) is the *SP* algorithm with the practical enhancement (complexity improvement) only and SP-both is the one with all enhancements. Besides, we study 4 approximate algorithms, Split, Merge, Greedy and Intersect. The first three are the adaptations of the existing trajectory simplification methods and the forth is proposed in this thesis.

All algorithms were implemented in C/C++ and run on a Linux platform with a 2.66GHz machine and 4GB RAM.

## 4.6.2 Relevance to Existing Studies

In this section, we conducted experiments to show how DPTS is relevant to existing studies.

<sup>&</sup>lt;sup>3</sup>http://research.microsoft.com/en-us/downloads/b16d359d-d164-469e-9fd4-daa38f2b2e13/

<sup>&</sup>lt;sup>4</sup>http://research.microsoft.com/apps/pubs/?id=152883

### **Bounds of DPTS wrt Existing Measurements**

In this part, we verify the theoretical bounds of the length (speed) error and the position error of DPTS as introduced in Section 4.3.1.

We vary the tolerance  $\epsilon_t$  on  $\{0.2, 0.4, 0.6, 0.8, 1\}$ . The results about length (speed) errors are shown in Figure 4.8(a), where the "length (speed) ratio" is defined to be  $\min_{i \in [1,n)} \{len(p'_i, p'_{i+1}|T')/len(p_i, p_{i+1}|T)\}$  where  $p'_i$  is the estimated position of  $p_i$  on T' for  $i \in [1,n]$ . Thus, the larger this ratio is, the more accurate the length (speed) information of the simplified trajectory is. Note that the length ratio and the speed ratio (with respect to a segment in the original trajectory) are exactly the same since the speed is equal the length divided by the time difference between the time stamps of the two end-points of the segment, and the time difference in the original trajectory is kept to be the same as the time difference in the simplified trajectory. We observe that the theoretical bound of the length (speed) ratio is usually good (e.g., it is about 0.92 when  $\epsilon_t = 0.4$ ).

The results about the position error are shown in Figure 4.8(b). We observe that the empirical position error is usually significantly smaller than the theoretical bound (by near to one order of magnitude). Besides, when  $\epsilon_t$  increases, the increase in the position error of DPTS becomes smaller. When  $\epsilon_t$  becomes large, the position error of DPTS keeps quite stable.

### **DPTS vs. PPTS**

In this part, we want to compare DPTS with <u>Position-Preserving Trajectory</u> <u>Simplification</u> (PPTS) in terms of two measurements, namely the position error and the direction error. In Section 4.3, we show that the position error of DPTS is *bounded* (Lemma 4.3.2) while the direction error of PPTS is *un-bounded* (Lemma 4.3.3). We study *how worst the position error of DPTS compared with PPTS is* and *how worst the* 

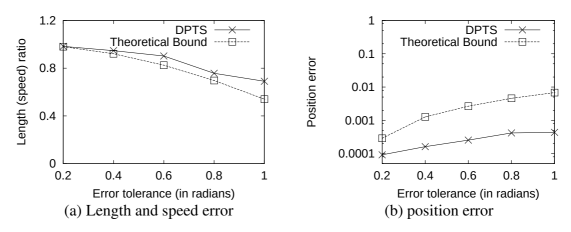


Figure 4.8: Verification of theoretical error bounds (Geolife, DPTS)

direction error of PPTS compared with DPTS is.

We adopt the *Douglas-Peucker* algorithm [24] for PPTS which is the most popular existing algorithm for PPTS [50, 9, 33], and we use our SP algorithm for DPTS. We vary  $\epsilon_t$  for DPTS. For a fair comparison, we enforce that the simplified trajectories from DPTS and PPTS have the same size. The results are shown in Figure 4.9(a) for position errors and in Figure 4.9(b) for direction errors. Consider Figure 4.9(a). It could be noticed that though the position errors of DPTS are usually larger than those of PPTS, the difference is small. For example, the ratio of the position errors is between 1.85 to 3. Consider Figure 4.9(b). We observe that the direction errors of PPTS are significantly larger than that of DPTS. For example, when  $\epsilon_t = 0.2$ , the ratio is more than 10. Besides, the direction errors of PPTS are greater than 2, a value greater than  $\pi/2$ , even with a small value of  $\epsilon_t$  and is nearly to  $\pi$ , the greatest possible direction error, with a medium value of  $\epsilon_t$ , which implies that PPTS can hardly preserves the direction information. In conclusion, our DPTS preserves the direction information by its nature and also the position information to a certain degree, but PPTS preserves the position information only but not the direction information.

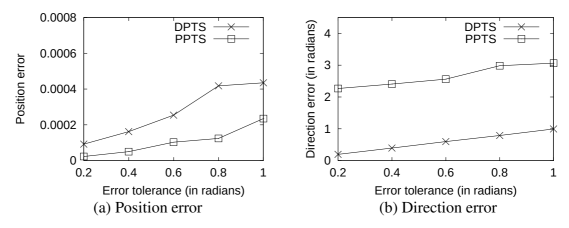


Figure 4.9: Comparison with existing PPTS (Geolife, DPTS)

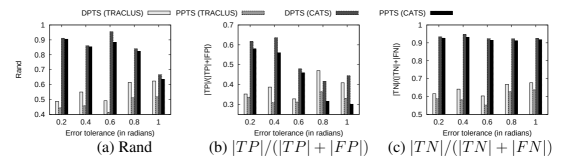


Figure 4.10: Trajectory Clustering Study (the Deer dataset, DPTS)

### **An Application Study (Trajectory Clustering)**

In Section 4.6.2, we compared DPTS and PPTS with their *favor-metrics* (i.e., the position error, favoring PPTS, and the direction error, favoring DPTS). In this section, we compare DPTS with PPTS with a *neutral* metric, the clustering quality, for a real-life application, trajectory clustering.

The main idea is as follows. Let  $\mathcal{D}$  be a set of raw trajectories. We perform DPTS (PPTS) on each trajectory in  $\mathcal{D}$  and obtain a set of simplified trajectories, denoted by  $\mathcal{D}^d$  ( $\mathcal{D}^p$ ). Then, we perform a clustering procedure on each of these two sets of trajectories and obtain the corresponding clustering results. We regard the clustering results based on  $\mathcal{D}$  as ground truth and measure the qualities of the clustering results on  $\mathcal{D}^d$  and  $\mathcal{D}^p$ . We verify DPTS by showing that the clustering results on  $\mathcal{D}^d$  are consistently better (closer to the ground truth) than those on  $\mathcal{D}^p$ .

Consider the clustering procedure on  $\mathcal{D}$  first. The clustering results based on  $\mathcal{D}$  (i.e., the cluster membership of each trajectory) could be represented by a binary matrix  $M_{n_t \times n_c}$ , where  $n_t$  is the number of trajectories in the set and  $n_c$  is the number of resulting clusters. Note that under some clustering mechanisms such as the one in [46], a trajectory could belong to multiple clusters. For each trajectory  $T \in \mathcal{D}$ , its cluster membership could be represented by an  $n_c$ -dimensional binary vector. For each pair of trajectories  $T_1$  and  $T_2$  in  $\mathcal{D}$ , we measure the similarity between  $T_1$  and  $T_2$  by the Euclidean distance between  $T_1$ 's cluster membership (which is a vector) and  $T_2$ 's cluster membership (which is a vector). If the distance is below a pre-set threshold  $\sigma$ , we regard  $T_1$  and  $T_2$  to be similar; otherwise, we regard  $T_1$  and  $T_2$  to be dissimilar. Thus, based on the clustering results on  $\mathcal{D}$ , we can always obtain a similarity matrix which indicates for each pair of two trajectories in  $\mathcal{D}$  whether they are similar or not. Let S be such a similarity matrix corresponding to  $\mathcal{D}$ . The distance threshold  $\sigma$  for deciding whether two trajectories are similar is set to 0.5 by default (all distances are normalized to [0,1]).

For a specific trajectory  $T \in \mathcal{D}$ , we denote its simplification in  $\mathcal{D}^d$  and  $\mathcal{D}^p$  by  $T^d$  and  $T^p$ , respectively.

Similarly, we perform the same clustering procedure on  $\mathcal{D}^d$  and  $\mathcal{D}^p$  as we did on  $\mathcal{D}$  and obtain their corresponding similarity matrices, denoted by  $S^d$  and  $S^p$ , respectively.

We measure the quality of the clustering on  $\mathcal{D}^d$  (and the clustering on  $\mathcal{D}^p$ ) as follows. For a pair of two trajectories  $T_1$  and  $T_2$  in  $\mathcal{D}$ , we have 4 cases. Case 1:  $T_1$  and  $T_2$  are similar (wrt S) and  $T_1^d$  and  $T_2^d$  (the simplified trajectories of  $T_1$  and  $T_2$  under DPTS) are similar (wrt  $S^d$ ). In this case, we have an occurrence of true positive (TP). Case 2:  $T_1$  and  $T_2$  are dissimilar (wrt S) and  $T_1^d$  and  $T_2^d$  are similar (wrt  $S^d$ ). In this case, we have an occurrence of false positive (FP). Similarly, Case 3 and Case 4 correspond to the occurrences of false negative (FN) and true negative (TN), respectively. We adopt

three measures in our experiments for measuring the clustering results. The first is called *Rand*, which is defined to be (|TP| + |TN|)/(|TP| + |FP| + |FN| + |TN|) where  $|\cdot|$  denotes the number of occurrences. The second and the third are defined to be |TP|/(|TP| + |FP|) and |TN|/(|TN| + |FN|), respectively. The larger the measure is, the better the clustering is.

For the trajectory clustering procedure, we adopt the *TRACLUS* algorithm [46] and the *CATS* algorithm [37]. According to [37], existing trajectory clustering algorithms fall in two categories. The first category includes those algorithms which take each trajectory *as a whole* for clustering while the second category includes the algorithms which use *sub-trajectories* for clustering. *CATS* is the state-of-the-art in the first category and *TRACLUS* is the state-of-the-art in the second category [37].

For the PPTS procedure, again, we adopt the popular *Douglas-Peucker* algorithm.

We vary the error tolerance  $\epsilon_t$  with the values of 0.2, 0.4, 0.6, 0.8 and 1 for DPTS. Figure 4.10(a), (b) and (c) show the results about the Rand measure, the measure of |TP|/(|TP| + |FP|) and the measure of |TN|/(|TN| + |FN|) on the Deer dataset, respectively.

We observe that the clustering based on the simplified trajectories returned by DPTS is consistently better than that based on the simplified trajectories returned by PPTS. This might be explained by the fact that the direction information is heavily used in the trajectory clustering algorithms and the direction information loss due to DPTS is bounded while that due to PPTS is un-bounded.

# **4.6.3** Performance Study of the Optimal Algs

In the part, we study the effects of 2 factors, namely the data size (i.e., |T|) and the error tolerance (i.e.,  $\epsilon_t$ ), on the performance of the optimal algorithms. We use 2 measures, namely the running time and the memory.

Effect of |T|. The values used for |T| are around 2k 4k, 6k, 8k and 10k ( $\epsilon_t$  is fixed to be 1). For each setting of |T|, we select a set of 10 trajectories each of which has its size near to this value and run DPTS on each of these trajectories. Then, we average the experimental results on these trajectories (this policy is used throughout our experiments without specification). Figure 4.11 show the results on Geolife. According to these results, SP-both is the fastest while DP is the slowest due to its high time complexity. Besides, the complexity improvement helps to reduce the running time dramatically (e.g., SP-theo is faster than SP by 2-3 orders of magnitude). This could be easily explained by the fact that with the complexity improvement, the cost of checking whether  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$  is reduced from O(n) to O(C) (C is a small constant). Though the practical enhancement improves the time efficiency a little, it helps to reduce the memory significantly (e.g., the memory occupied by SP-theo is 1-3 orders of magnitude larger than that occupied by SP-both and the difference increases when |T| increases on Geolife).

The experimental results on T-Drive are similar and thus they are omitted.

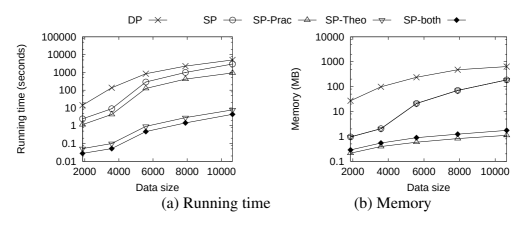


Figure 4.11: Effect of data size |T| (Geolife, DPTS)

Effect of  $\epsilon_t$ . The values used for  $\epsilon_t$  are 0.2, 0.4, 0.6, 0.8 and 1 in radians (|T| is fixed to be 5k). Figure 4.12 shows the results on Geolife. According to these results,  $\epsilon_t$  affects the SP algorithms only. Specifically, the running times of all SP algorithms increase slightly when  $\epsilon_t$  becomes larger. This is because a larger  $\epsilon_t$  usually results in  $G_{\epsilon_t}$  with

more edges and thus the BFS process on  $G_{\epsilon_t}$  needs more time.

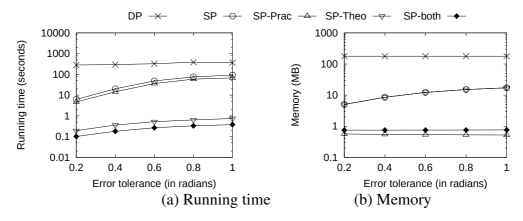


Figure 4.12: Effect of error tolerance  $\epsilon_t$  (Geolife, DPTS)

Compression Rate. We also study the effect of  $\epsilon_t$  on the *size ratio* which is defined to be equal to  $\sum_{T' \in \mathcal{D}'} |T'| / \sum_{T \in \mathcal{D}} |T|$ , where  $\mathcal{D}$  is the set of raw trajectories and  $\mathcal{D}'$  is the set of the corresponding simplified trajectories. Note that a smaller size ratio means a higher compression rate. The results are shown in Figure 4.13. We have the following observations. First, the size ratio decreases significantly when we increase the tolerance from 0 slightly. This is good since it implies that under DPTS, the trajectory data could be simplified significantly with a small error. Second, we observe that the size ratio is strictly smaller than 1 (e.g., it is about 0.9 for the Geolife datasets) even if the error tolerance is set to be 0. This implies that the real-life trajectories usually involve a certain degree of redundancy and could be simplified without incurring any error.

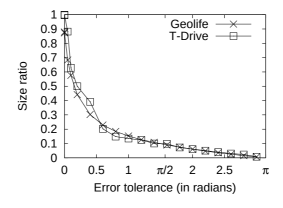


Figure 4.13: Compression rate, DPTS

Scalability Test. Figure 4.14 shows the results of the scalability test on the optimal algorithms. We only show the results of SP-theo and SP-both since the other optimal algorithms are not scalable on large datasets due to their expensive time complexities. According to these results, both SP-theo and SP-both are scalable to large trajectory datasets with millions of positions, and SP-both runs slightly faster than SP-theo. It is noted that SP-both occupies significantly less memory than SP-theo and thus SP-both is more scalable than SP-theo. This is because SP-both does not materialize  $G_{\epsilon_t}$  explicitly while SP-theo does.

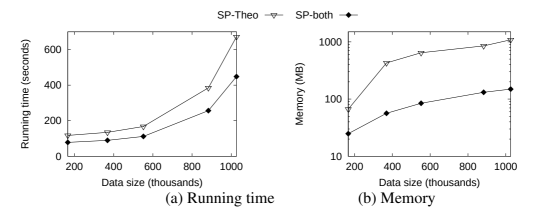


Figure 4.14: Scalability test for exact algorithms (Geolife, DPTS)

#### 4.6.4 Performance Study of the Approx. Algs

In this part, we study the effects of |T| and  $\epsilon_t$  on the approximate algorithms. We use 3 measures, namely the running time, the memory and the approximation error. The approximation error of an approximate algorithm is defined to be  $|T'|/|T^*|$ , where T' is the simplified trajectory returned by the approximate algorithm on a given raw trajectory and  $T^*$  is the simplified trajectory returned by an optimal algorithm on the same raw trajectory.

**Effect of** |T|. The values used for |T| are around 200k, 400k, 600k, 800k and 1000k ( $\epsilon_t$  is fixed to be 1). Figure 4.15 shows the results. According to these results, Intersect is the fastest, which is at least 1 order of magnitude faster than other approximate

algorithms. This is because Intersect runs in linear time while the other algorithms run in quadratic time in the worst case. Besides, Intersect occupies the least memory and Greedy occupies slightly more memory than Intersect (though the difference in Figure 4.15(b) is not obvious).

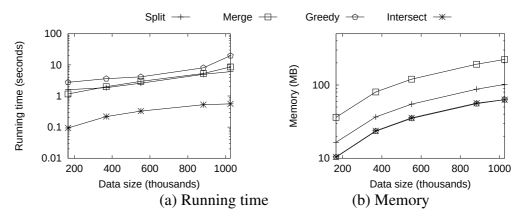


Figure 4.15: Effect of data size |T| (Geolife, DPTS)

Effects of  $\epsilon_t$ . Figure 4.16 shows the effects of  $\epsilon_t$  on the approximate algorithms, where we vary  $\epsilon_t$  with 0.2, 0.4, 0.6, 0.8 and 1 (|T| is fixed to be 500k). According to these results, Greedy runs faster than Split and Merge with small  $\epsilon_t$ 's. This is because with a smaller  $\epsilon_t$ , it is less likely that a long sequence of consecutive segments could be approximated with one segment and thus the cost of checking the error of the segment linking the start position and the end position is small.

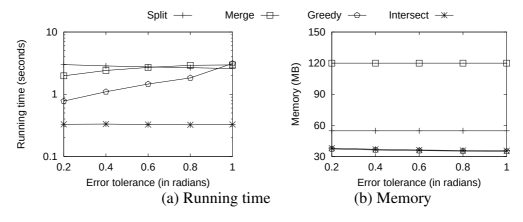


Figure 4.16: Effect of the error tolerance  $\epsilon_t$  (Geolife, DPTS)

**Compression Rate & Approximation Error.** Figure 4.17(a) shows the results on the compression rates of the approximate algorithms. We also show the theoretical bound

of the size the trajectory returned by Intersect (Lemma 4.5.3). Figure 4.17(b) shows the results on the approximate errors of the approximate algorithms. Both these results verify our Intersect algorithm.

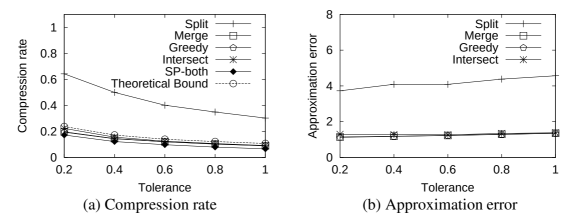


Figure 4.17: Compression rates and approximate errors of the approximate algoritms (Geolife, DPTS)

**Scalability Test.** The largest trajectory in our real datasets contains around 2M positions only. In order to generate larger trajectories, we concatenate multiple trajectories in ascending order of their time stamps into one. Figure 4.18 shows the results of the scalability test on the approximate algorithms. According to these results, Intersect is very fast on large datasets with more than 20M positions. For example, Intersect runs in 13.3s on the trajectory with 24,876,978 positions. In contrast, the running times of other approximate algorithms increase much faster when |T| increases.

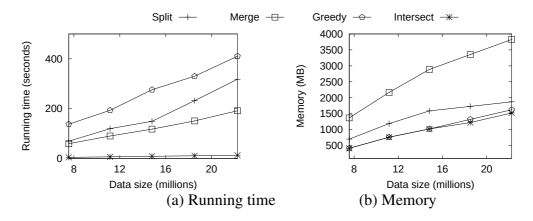


Figure 4.18: Scalability test for approximate algorithms (Geolife, DPTS)

### 4.7 Conclusion

In this chapter, we propose direction-preserving trajectory simplification, which has not been studied in the literature, as a novel alternative to the traditional position-preserving trajectory simplification. We propose an optimal algorithm called *SP* and an approximate algorithm called *Intersect*. We conducted experiments to show the efficiency and the scalability of our proposed methods.

### CHAPTER 5

### CONCLUSION AND FUTURE WORK

In this thesis, we introduce three techniques for boosting the spatial computations which are central to some location-based services, namely the collective spatial keyword query, worst-case optimized spatial matching, and direction-preserving trajectory simplification.

Motivated by the fact that DPTS requires users to specify an error tolerance which users might not know how to set properly in some cases (e.g., the error tolerance could only be known at some future time and simply setting one error tolerance does not meet the needs since the simplified trajectories would usually be used in many different applications which accept different error tolerances), I plan to study the problem of minimizing the error while achieving a pre-defined simplification size for trajectory simplification.

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#### APPENDIX A

### PROOFS OF LEMMAS/THEOREMS

# A.1 The Approximation Factor of MaxSum-Appro in General Case

In this part, we show that the approximation factor of MaxSum-Appro with a general setting of  $\alpha$  is equal to  $(2 - \sqrt{2}/2 \cdot \alpha)$ .

We use the same notations as defined in the proof of Theorem 2.3.2.

Consider  $cost(S_o)$ . Same as the proof of Theorem 2.3.2, we have  $\max_{o' \in S_o} d(o',q) = d(o,q) = r_2$  and  $\max_{o_1,o_2 \in S_o} d(o_1,o_2) \geq d(o,o_f) = r_1$ . As a result, we have

$$cost(S_o) \ge \alpha \cdot r_2 + (1 - \alpha) \cdot r_1 \tag{A.1}$$

Consider cost(S'). Same as the proof of Theorem 2.3.2, we consider two cases.

Case 1:  $r_1 \leq \sqrt{2}r_2$  (See Figure 2.2(a) for illustration). In this case, we have  $\max_{o_1,o_2\in S'}d(o_1,o_2)\leq d(a,b)=2\sqrt{r_1^2-r_1^4/4r_2^2}$ . Recall that  $\max_{o'\in S'}d(o',q)=d(o,q)=r_2$ . As a result, we have

$$cost(S') \le \alpha \cdot r_2 + (1 - \alpha) \cdot 2\sqrt{r_1^2 - r_1^4/4r_2^2}$$
 (A.2)

Therefore,

$$\frac{\cos t(S')}{\cos t(S_o)} \leq \frac{\alpha \cdot r_2 + (1 - \alpha) \cdot 2\sqrt{r_1^2 - r_1^4/4r_2^2}}{\alpha \cdot r_2 + (1 - \alpha) \cdot r_1} 
= 1 + \frac{(1 - \alpha) \cdot (2\sqrt{r_1^2 - r_1^4/4r_2^2} - r_1)}{\alpha \cdot r_2 + (1 - \alpha) \cdot r_1} 
= 1 + \frac{(1 - \alpha) \cdot (2\sqrt{1 - r_1^2/4r_2^2} - 1)}{\alpha \cdot r_2/r_1 + 1 - \alpha} 
< 1 + \frac{(1 - \alpha) \cdot (2\sqrt{1 - 0} - 1)}{\alpha \cdot \sqrt{2}/2 + 1 - \alpha} 
= 1 + \frac{1 - \alpha}{\alpha \cdot \sqrt{2}/2 + 1 - \alpha} 
= 2 - \frac{\sqrt{2}/2 \cdot \alpha}{1 - (1 - \sqrt{2}/2) \cdot \alpha} 
< 2 - \frac{\sqrt{2}/2 \cdot \alpha}{1 - 0} = 2 - \sqrt{2}/2 \cdot \alpha$$

Case 2:  $r_1 > \sqrt{2}r_2$  (See Figure 2.2(b) for illustration). Similar to Case 1, it could be verified that  $\max_{o_1,o_2\in S'}d(o_1,o_2)\leq d(a,b)=2r_2$ . As a result, we have

$$cost(S') \le \alpha \cdot r_2 + (1 - \alpha) \cdot 2r_2 \tag{A.3}$$

Therefore,

$$\frac{cost(S')}{cost(S_o)} \leq \frac{\alpha \cdot r_2 + (1-\alpha) \cdot 2 \cdot r_2}{\alpha \cdot r_2 + (1-\alpha) \cdot r_1} = \frac{\alpha + (1-\alpha) \cdot 2}{\alpha + (1-\alpha) \cdot r_1/r_2}$$

$$\leq \frac{2-\alpha}{\alpha + (1-\alpha) \cdot \sqrt{2}} < 2-\alpha < 2-\sqrt{2}/2 \cdot \alpha$$

Thus, by combining Case 1 and Case 2, we have  $cost(S') \leq (2 - \sqrt{2}/2 \cdot \alpha) \cdot cost(S_o)$ , which finishes the discussion.

# A.2 Theoretical Error Bounds wrt the Synchronous Euclidean Distance

In this part, we show that when the mapping function  $M_{\mathcal{S}}(\cdot, \cdot)$  (i.e., Synchronous Euclidean Distance) is used, the position error of DPTS is bounded (Lemma A.2.1) and the direction error of PPTS is un-bounded (Lemma A.2.2).

**Lemma A.2.1 (Bounded Position Error)** Let T be a trajectory and T' be an  $\epsilon_t$ simplification of T with  $\epsilon_t < \pi/2$ . For each position  $p_i$  in T where  $i \in [1, n]$ , we have

$$d(p_i, p_i') \le 0.5 \cdot (1 + 1/\cos(\epsilon_t)) \cdot L_{max}$$

where 
$$p'_{i} = M_{\mathcal{C}}(p_{i}, T')$$
 and  $L_{max} = \max_{i \in [1,n]} len(p_{s_{i}}, p_{s_{i+1}} | T')$ .

**Proof.** Let  $\overline{p_{s_k}p_{s_{k+1}}}$  be the segment of T' such that  $p_{s_k}$  is the last position with  $s_k \leq i$  and  $p_{s_{k+1}}$  is the first position with  $s_{k+1} \geq i+1$ . Consider Figure A.1(a) for illustration.

By using triangle inequality, we have

$$d(p_i, p_i') \le len(p_{s_k}, p_i|T) + d(p_{s_k}, p_i')$$

and

$$d(p_i, p'_i) \le len(p_i, p_{s_{k+1}}|T) + d(p'_i, p_{s_{k+1}})$$

By using the above two equations, we have

$$\begin{split} d(p_i, p_i') &\leq len(p_{s_k}, p_i|T) + d(p_{s_k}, p_i') \\ &+ len(p_i, p_{s_{k+1}}|T) + d(p_i', p_{s_{k+1}}) \\ &= len(p_{s_k}, p_{s_{k+1}}|T) + len(p_{s_k}, p_{s_{k+1}}|T') \\ &\leq 1/\cos(\epsilon_t) \cdot len(p_{s_k}, p_{s_{k+1}}|T') + len(p_{s_k}, p_{s_{k+1}}|T') \\ &= (1 + 1/\cos(\epsilon_t)) \cdot len(p_{s_k}, p_{s_{k+1}}|T') \\ &\leq (1 + 1/\cos(\epsilon_t)) \cdot L_{max} \end{split}$$

which completes the proof.

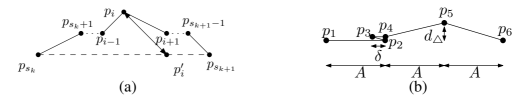


Figure A.1: Proofs of Lemma A.2.1 and Lemma A.2.2

**Lemma A.2.2 (Unbounded Direction Error)** Let T be a trajectory and T' be a (direction-based)  $\epsilon_t$ -simplification of T with  $\epsilon_t < \pi/2$ . Let  $T_C$  be a (position-based) simplified trajectory of T such that  $|T_C| = |T'|$  and the error of  $T_C$  under the Synchronous Euclidean Distance is minimized. There exists a dataset such that  $\epsilon(T_C) \approx \pi$  and  $\epsilon(T') \approx 0$ .

**Proof.** We prove by constructing a problem instance as shown in Figure A.1(b).  $T=(p_1,p_2,p_3,p_4,p_5,p_6)$  is a trajectory, where all positions except  $p_5$  are along a horizontal line such that  $d(p_2,p_1)=d(p_4,p_1)=A$ ,  $d(p_3,p_1)=A-\delta$  and  $d(p_6,p_1)=3A$ . Here, A is a positive real number. Besides,  $p_5$  is located a little bit above the horizontal line with its perpendicular distance to the horizontal line equal to  $d_{\triangle}$ . We have  $\delta << d_{\triangle}$ .

Suppose that we can only keep 5 positions in the simplified trajectory. In other words, we have to remove 1 position from the 6 positions. If we consider preserving

the direction information,  $p_4$  will be removed and thus  $T'=(p_1,p_2,p_3,p_5,p_6)$ . As a result,  $\epsilon(T')\approx 0$ . If we consider preserving the position information,  $p_3$  will be removed (since the position error of  $p_3$  under the Synchronous Euclidean Distance is the smallest) and thus  $T_{\mathcal{C}}=(p_1,p_2,p_4,p_5,p_6)$ . As a result,  $\epsilon(T_{\mathcal{C}})\approx \pi$ .

## A.3 Adaptations of Existing Trajectory Simplification Methods

The existing trajectory simplification methods described in Section 4.3 can be adapted to the DPTS problem. However, these adaptations have limitations. We discuss both the adaptations and their limitations next.

Let  $T=(p_1,p_2,...,p_n)$  be a trajectory and  $\epsilon_t$  be the error tolerance.

**Split.** It splits T at a *splitting position* into two sub-trajectories if the error of approximating T with one segment exceeds  $\epsilon_t$ . Let i be  $\arg\max_{1\leq h\leq n}\{\triangle(\theta(\overline{p_hp_{h+1}}),\theta(\overline{p_1p_n}))\}$ . We select  $p_i$  as the splitting position. It then performs the same procedure recursively on each of the sub-trajectories. At the end, it approximates each resulting trajectory with one segment.

The time complexity of Split is  $O(n^2)$  because in the worst case, it always chooses the second position of the remaining trajectory as the splitting point.

Merge. It regards each segment in T as a sub-trajectory. It then iteratively merge two adjacent sub-trajectories  $T_1$  and  $T_2$  into one if the error of approximating the merged trajectory with one segment is bounded by  $\epsilon_t$  until no such merge operations are possible. When multiple candidates of two adjacent sub-trajectories for merging are available, we choose the one with the smallest error when the corresponding merged trajectory is approximated with one segment. At the end, it approximates each resulting sub-trajectory with one segment.

The time complexity of Merge is  $O(n^2)$  since, in the worst case, it always merges the first two remaining sub-trajectories and the final simplified returned is  $\overline{p_1p_n}$ .

**Greedy.** Let T' be the simplified trajectory to be computed by Greedy, which is initialized as  $(p_1)$  at the beginning. Greedy scans the positions of T starting from  $p_1$  sequentially. It iteratively finds the position  $p_i$  such that  $\epsilon(\overline{p_ep_i}) \leq \epsilon_t$  and  $\epsilon(\overline{p_ep_{i+1}}) > \epsilon_t$ , where  $p_e$  is the last position in T' currently maintained in the algorithm, and then appends  $p_i$  to T'. At the end, it appends  $p_n$  to T'.

The time complexity of Greedy is  $O(n^2)$ . This is because, in the worst case, where the output T' contains only two positions,  $p_1$  and  $p_n$ , it has to compute  $\epsilon(\overline{p_1p_3})$ ,  $\epsilon(\overline{p_1p_4})$ , ...,  $\epsilon(\overline{p_1p_n})$ . Since each  $\epsilon(\overline{p_1p_3})$  takes O(n) time, the time complexity is  $O(n^2)$ .

### A.4 The DP Method

In the following, we design a <u>dynamic programming</u> (DP) algorithm for the DPTS problem. Let T[i,j] be the sub-trajectory of T, which starts at  $p_i$  and ends at  $p_j$  where  $1 \le i < j \le n$ , i.e.,  $T[i,j] = (p_i, p_{i+1}, ..., p_j)$ . We denote by S[i,j] the optimal solution of the DPTS problem where the trajectory input is T[i,j] and the error tolerance is still  $\epsilon_t$ . Let N[i,j] denote the number of segments maintained in S[i,j] (which is equal to the size of the optimal solution (i.e., S[i,j]) minus one). Note that minimizing the number of segments in the simplified trajectory is exactly equal to minimizing the size of the simplified trajectory.

Consider S[i, j]. There are two cases.

- Case 1:  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$ . That is, T[i,j] could be approximated by one segment (i.e.,  $\overline{p_ip_j}$ ) with the error bounded by the tolerance. In this case, we have  $S[i,j] = (p_i, p_j)$  and N[i,j] = 1.
- ullet Case 2:  $\epsilon(\overline{p_ip_j}) > \epsilon_t$ . In this case, T[i,j] cannot be approximated with one

segment only such that the error tolerance constraint is satisfied. Thus, there exists at least one position  $p_{k^*}$  in T[i,j] excluding its start position and its end position where  $i < k^* < j$ . Besides, we have  $N[i,j] = N[i,k^*] + N[k^*,j]$  and  $k^* = \arg\min_{i < k < j} \{N[i,k] + N[k,j]\}$ .

In conclusion, we can compute N[i,j] in the way as shown in Equation (A.4).

$$N[i,j] = \begin{cases} 1 & \text{if } \epsilon(\overline{p_i p_j}) \le \epsilon_t \\ \min_{i < k < j} \{N[i,k] + N[k,j]\} & Otherwise \end{cases}$$
(A.4)

Based on Equation (A.4), we can design a DP algorithm for the DPTS problem. Note that N[1, n] corresponds to the number of segments maintained in the optimal solution of the DPTS problem with the input as T.

Complexity Analysis. For each problem instance N[i,j], we first check whether  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$ . The cost is simply O(n) since it computes  $\Delta(\overline{p_ip_j},\overline{p_hp_{h+1}})$  for  $i \leq h < j$  in order to obtain  $\epsilon(\overline{p_ip_j})$ . Besides, it takes O(1) and O(n) time to obtain N[i,j] in Case 1 and Case 2, respectively. Thus, the overall cost for a specific problem instance of N[i,j] is still O(n). Since we have in total  $O(n^2)$  possible problem instances of N[i,j], the time complexity of the DP algorithm is  $O(n^2 \cdot n) = O(n^3)$ .

### A.5 Complete Proof of Lemma 7

Since  $\overline{p_ip_j}$  is  $\frac{\epsilon_t}{2}$ -feasible,  $fdr(T[i,j]|\epsilon_t/2)$  is non-empty.

We prove with two steps.

First, we deduce that for any two segments between i and j in T,  $\overline{v_hv_{h+1}}$  and  $\overline{v_{h'}v_{h'+1}}$  ( $i \leq h < h' < j$ ), we have  $\triangle(\theta(\overline{p_hp_{h+1}}), \theta(\overline{p_{h'}p_{h'+1}})) \leq \epsilon_t$ . Let  $\theta_1 = \theta(\overline{p_hp_{h+1}})$  and  $\theta_2 = \theta(\overline{p_{h'}p_{h'+1}})$ . Since  $fdr(T[i,j]|\epsilon_t/2)$  is non-empty, we know  $fdr(\overline{p_hp_{h+1}}|\epsilon_t/2)$  and  $fdr(\overline{p_{h'}p_{h'+1}}|\epsilon_t/2)$  intersects. Let  $\theta_3$  be a direction that

is in both  $fdr(\overline{p_hp_{h+1}}|\epsilon_t/2)$  and  $fdr(\overline{p_{h'}p_{h'+1}}|\epsilon_t/2)$ . Thus,  $\triangle(\theta_1,\theta_3) \leq \epsilon_t/2$  and  $\triangle(\theta_2,\theta_3) \leq \epsilon_t/2$ . As a result, according to Equation (4.1), we have

$$\Delta(\theta_1, \theta_3) = \min\{|\theta_1 - \theta_3|, 2\pi - |\theta_1 - \theta_3|\} \le \epsilon_t/2 \tag{A.5}$$

$$\Delta(\theta_2, \theta_3) = \min\{|\theta_2 - \theta_3|, 2\pi - |\theta_2 - \theta_3|\} < \epsilon_t/2 \tag{A.6}$$

Next, we show that  $\triangle(\theta_1, \theta_2) \le \epsilon_t$ . Consider four cases.

Case 1:  $|\theta_1 - \theta_3| \le 2\pi - |\theta_1 - \theta_3|$  and  $|\theta_2 - \theta_3| \le 2\pi - |\theta_2 - \theta_3|$ . In this case, we have  $|\theta_1 - \theta_3| \le \epsilon_t/2$  and  $|\theta_2 - \theta_3| \le \epsilon_t/2$ . Thus,

$$|\theta_1 - \theta_2| = |\theta_1 - \theta_3 + \theta_3 - \theta_2| \le |\theta_1 - \theta_3| + |\theta_2 - \theta_3| \le \epsilon_t$$

Therefore,  $\triangle(\theta_1, \theta_2) = \min\{|\theta_1 - \theta_2|, 2\pi - |\theta_1 - \theta_2|\} \le \epsilon_t$ .

Case 2:  $|\theta_1 - \theta_3| > 2\pi - |\theta_1 - \theta_3|$  and  $|\theta_2 - \theta_3| \le 2\pi - |\theta_2 - \theta_3|$ . In this case, we have  $|\theta_1 - \theta_3| > 2\pi - \epsilon_t/2$  and  $|\theta_2 - \theta_3| \le \epsilon_t/2$ .

$$|\theta_1 - \theta_2| = |\theta_1 - \theta_3 - (\theta_2 - \theta_3)| \ge |\theta_1 - \theta_3| - |\theta_2 - \theta_3| >$$

$$2\pi - \epsilon_t/2 - \epsilon_t/2 = 2\pi - \epsilon_t$$

Therefore,  $\triangle(\theta_1, \theta_2) = \min\{|\theta_1 - \theta_2|, 2\pi - |\theta_1 - \theta_2|\} \le \epsilon_t$ .

Case 3:  $|\theta_1 - \theta_3| \le 2\pi - |\theta_1 - \theta_3|$  and  $|\theta_2 - \theta_3| > 2\pi - |\theta_2 - \theta_3|$ . This case is symmetric with Case 2 and the proof is similar.

Case 4:  $|\theta_1 - \theta_3| > 2\pi - |\theta_1 - \theta_3|$  and  $|\theta_2 - \theta_3| > 2\pi - |\theta_2 - \theta_3|$ . In this case, we have  $2\pi - |\theta_1 - \theta_3| \le \epsilon_t/2$  and  $2\pi - |\theta_2 - \theta_3| \le \epsilon_t/2$ . As a result, we have  $|\theta_1 - \theta_3| > \pi$  which implies that  $\theta_1$  and  $\theta_3$  are at different sides of the x-axis, and  $|\theta_2 - \theta_3| > \pi$  which implies that  $\theta_2$  and  $\theta_3$  are at different sides of the x-axis as well. Without loss of generality, assume that  $\theta_3$  is at the upper side of the x-axis. Then, both  $\theta_1$  and  $\theta_2$  are at the lower side. Furthermore, we assume assume  $\theta_1 \ge \theta_2$  without loss

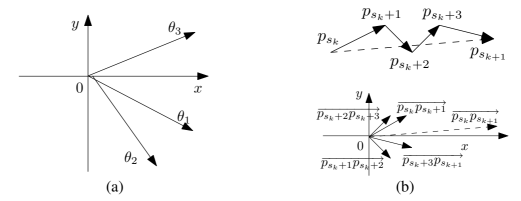


Figure A.2: Proof of Lemma 4.5.2

of generality. Then,  $|\theta_1 - \theta_2| \le 2\pi - |\theta_2 - \theta_3| \le \epsilon_t/2 < \epsilon_t$ . Consider Figure A.2(a) for illustration. Therefore,  $\triangle(\theta_1, \theta_2) \le |\theta_1 - \theta_2| \le \epsilon_t$ .

By combining the above four cases, we have  $\triangle(\theta_1, \theta_2) \le \epsilon_t$ .

Second, we show that  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$ . According to the results in the first step, for any two segments between  $p_i$  and  $p_j$  in T, the angular difference between their directions is at most  $\epsilon_t$ . It is easy to verify that there exists an angular range [a,b] such that the directions of all the segments between  $p_i$  and  $p_j$  in T fall in [a,b] and the span of [a,b] is bounded by  $\epsilon_t$  (where the definition of "span" can be found in the proof of Lemma 4.4.2). Given two positions p and p', we denote the *vector* from p to p' by  $\overrightarrow{pp'}$ . Besides, it could be verified that  $\theta(\overline{p_ip_j})$  falls in the angular range [a,b] by utilizing the fact that  $\overline{p_ip_j} = \sum_{h=i}^{j-1} \overline{p_hp_{h+1}}$ . For illustration, consider Figure A.2(b), where [a,b] corresponds to  $[\theta(\overline{p_{i+1}p_{i+2}}), \theta(\overline{p_{i+2}p_{i+3}})]$ . Therefore, we know  $\Delta(\theta(\overline{p_ip_j}), \theta(\overline{p_hp_{h+1}})) \leq \epsilon_t$  for  $i \leq h < j$  since the angular difference between any two directions falling in range [a,b] is smaller than that the angular difference between a and b, which is bounded by  $\epsilon_t$ . Thus, we know  $\epsilon(\overline{p_ip_j}) \leq \epsilon_t$ .

### A.6 Complete Proof of Lemma 9

Let  $T=(p_1,...,p_n)$ ,  $T'=(p_{s_1},...,p_{s_m})$  and  $T''=(p_{r_1},...,p_{r_l})$ . By definition, we have  $s_1=r_1=1$  and  $s_m=r_l=n$ . Note that n=|T|, m=|T'| and l=|T''|.

First, we prove that  $fdr(T[r_k,r_{k+1}]|\epsilon_t/2)$  is non-empty for  $1 \leq k < l$ . Consider a specific  $k \in [1,l-1]$ . Since T'' is an  $\epsilon_t/2$ -simplification, i.e.,  $\epsilon(T'') \leq \epsilon_t/2$ , we know  $\epsilon(\overline{p_{r_k}p_{r_{k+1}}}) \leq \epsilon_t/2$ . As a result, we know  $\Delta(\theta(\overline{p_hp_{h+1}}),\theta(\overline{p_{r_k}p_{r_{k+1}}})) \leq \epsilon_t/2$ , which further implies that  $\theta(\overline{p_{r_k}p_{r_{k+1}}})$  falls in  $fdr(\overline{p_hp_{h+1}}|\epsilon_t/2)$  for  $r_k \leq h < r_{k+1}$ . Thus,  $fdr(T[r_k,r_{k+1}]|\epsilon_t/2)$  is non-empty.

Next, we prove  $m \leq l$  by contradiction. Assume that m > l. In the following, we want to show that  $s_k \geq r_k$  for k = 1, 2, ..., l. When k = 1, it is true since  $s_1 = r_1 = 1$ . Consider k = 2. According to Algorithm *Intersect*, we know that  $fdr(T[s_1,s_2]|\epsilon_t/2)$  is non-empty but  $fdr(T[s_1,s_2+1]|\epsilon_t/2)$  is empty. We can say that  $s_2$  is the *greatest* possible position such that  $fdr(T[s_1, s_2]|\epsilon_t/2)$  is non-empty. Besides, since  $fdr(T[r_1, r_2]|\epsilon_t/2)$  is non-empty and  $r_1 = s_1$ , we derive that  $s_2 \geq r_2$ . Consider k=3. Since  $fdr(T[r_2,r_3]|\epsilon_t/2)$  is non-empty and  $s_2\geq r_2$ , we consider two cases. Case 1:  $s_2 < r_3$ . In this case, we know that  $fdr(T[s_2, r_3]|\epsilon_t/2)$ is non-empty and thus  $s_3 \geq r_3$ . Case 2:  $s_2 \geq r_3$ . In this case, we also deduce that  $s_3 \ge r_3$  (since  $s_3 > s_2$ ). By combining these two cases, we conclude that  $s_3 \geq r_3$ . Continuing the above procedure, we can deduce that  $s_k \geq r_k$  for k = 1, 2, ..., l. Since m > l, we have  $s_l < s_m$  and thus  $s_l + 1 \le s_m = r_l$ . Note that  $fdr(T[r_{l-1}, r_l] | \epsilon_t/2) = fdr(T[r_{l-1}, s_{l-1}] | \epsilon_t/2) \cap fdr(T[s_{l-1}, s_l+1] | \epsilon_t/2) \cap fdr(T[s_l+1] | \epsilon_t/2) \cap f$  $1, s_m ||\epsilon_t/2)$  (When  $s_l + 1 = s_m$ , we define  $fdr(T[s_l + 1, s_m]|\epsilon_t/2)$  to be the universe angular range). Besides, since  $fdr(T[r_{l-1},r_l]|\epsilon_t/2)$  is non-empty, we deduce that  $fdr(T[s_{l-1}, s_l + 1] | \epsilon_t/2)$  is non-empty, which, however leads to a contradiction because according to algorithm *Intersect*,  $fdr(T[s_{l-1}, s_l + 1] | \epsilon_t/2)$  is empty. Thus, we conclude that  $m \leq l$ , which implies that  $|T'| \leq |T''|$ .