Query Processing over Probabilistic Data with Gaussian Distributions

Tingting DONG
Abstract

The field of uncertain data management has received extensive attention of researchers due to the increasing demand for managing uncertain data in a large variety of real-world applications such as sensor networks, location-based services, monitoring and surveillance. Uncertainty can occur for different reasons, including measurement errors and noises in sensors, privacy-preserving transformations of sensitive records and the limited confidence in the output of predictive models.

Managing uncertainty involves modeling, representing, querying, and indexing uncertain data. Answering queries over uncertain databases poses more challenges than that over traditional databases because managing uncertainty usually means costly probability computations. Hence, it is crucial to develop efficient solutions when managing uncertain data. In this thesis, we model uncertainty probabilistically and represent each uncertain object in the database using a Gaussian distribution, which is a typical probability distribution widely used in statistics, pattern recognition, and machine learning. We consider the following three types of queries or searches over probabilistic data with Gaussian distributions.

First, we study the probabilistic range query, which is an important query in the field of uncertain data management. A probabilistic range query returns objects in the database that exist within a specified range from the query object with probabilities no less than a given probability threshold. The query object can be either a certain point or an uncertain object represented by a Gaussian distribution.

We propose several effective filtering techniques by analyzing the properties of Gaussian distribution. The proposed filtering techniques can significantly reduce the number of objects that need to be verified by expensive probability computation. In this way, we can avoid unnecessary computations and hence save cost. To support efficient query processing, we further propose a novel indexing method to enable filtering unpromising objects by groups rather than individually. We develop the indexing method by extending the existing R-tree and improve it based on our analysis of Gaussian distribution. This indexing method can effectively organize objects and greatly enhance the performance of query processing. Extensive experiments on real datasets demonstrate the efficiency and effectiveness of our proposed approach.

Second, we investigate the nearest neighbor search. As one of the commonest queries over location information, the distance-based nearest neighbor search, which finds closest
objects to a given query point, has extensive applications in many areas. There have been considerable efforts to extend nearest neighbor search over traditional location information to uncertain location information. An example is the expected distance, which defines the distance over uncertain location information. Following this trend, we represent uncertain locations using Gaussian distributions and assume that the closeness between each Gaussian object and the query point is measured by their expected distance. Under this setting, we consider the problem of \textit{k-expected nearest neighbor search} over Gaussian objects. The result objects are ones that have the top-\(k\) smallest expected distances to the query point.

We analyze properties of expected distance on Gaussian distribution mathematically and derive the lower bound and upper bound of the distance. Based on our analysis, we propose three novel approaches to efficiently solve this problem. The proposed approaches can prune unpromising objects whose lower bound distances are larger than upper bound or expected distances of candidate objects without computing their actual expected distances. We only compute exact expected distances for candidate objects and finally return the top-\(k\) smallest ones. To further improve the performance, we utilize R-tree to index objects and their lower bound distances and upper bound distances. The proposed approaches can effectively reduce the number of exact distance computation which is rather expensive. The efficiency and effectiveness of our approaches are demonstrated through extensive experiments.

Finally, we explore the problem of similarity search, which is a crucial task in many real-world applications such as multimedia databases, data mining, and bioinformatics. In this work, we investigate similarity search on uncertain data represented by Gaussian distributions. The query object is also represented by a Gaussian distribution. By employing \textit{Kullback-Leibler divergence} (KL-divergence) to measure the similarity between two Gaussian distributions, our goal is to search a database for the top-\(k\) Gaussian distributions similar to a given query Gaussian distribution. Especially, we consider non-correlated Gaussian distributions, where there are no correlations between dimensions and their covariance matrices are diagonal.

To support query processing, we propose two types of novel approaches utilizing the notions of \textit{rank aggregation} and \textit{skyline queries}. The first type presorts all objects in the database on their attributes and computes result objects by merging candidates from each presorted list. The second one transforms the problem to the computation of \textit{dynamic skyline queries}. We extend and modify the branch-and-bound skyline (BBS) algorithm, which is proposed to answer skyline queries, and develop a novel algorithm to solve this
problem. We demonstrate the efficiency and effectiveness of our approaches through a comprehensive experimental performance study.

In general, we provide a comprehensive view of managing probabilistic data with Gaussian distributions. We believe that our contributions are multi-dimensional and will be extended over time. First of all, we think our mathematical analyses of probabilistic range query, nearest neighbor search, and similarity search over Gaussian objects will benefit not only the existing related applications, but also potential studies of other applications over data represented by Gaussian distributions. Furthermore, our proposed non-trivial algorithms and indexing structures for efficient query processing, will not only enrich user experience by speed in the real world, but also provide valuable insights and references for developing solutions to other problems. Last but not least, we have conducted extensive experimental evaluations on the performance of our proposed solutions.
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<td>Bounding Box</td>
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<td>BBS</td>
<td>Branch-and-Bound Skyline</td>
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<td>ESQED</td>
<td>Expected SQuated Euclidean Distance</td>
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<td>LB</td>
<td>Lower Bound</td>
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<td>MBB</td>
<td>Minimum Bounding Box</td>
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<td>MBR</td>
<td>Minimum Bounding Rectangle</td>
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<td>PRQ</td>
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<td>SQED</td>
<td>SQuated Euclidean Distance</td>
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<td>UB</td>
<td>Upper Bound</td>
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Chapter 1

Introduction

1.1 Research Background

In recent years, the rapid advancement in data acquiring devices has led to the increasing availability of massive data in a wide variety of fields such as information retrieval, data integration, mobile robotics, sensor networks, location-based services, monitoring and surveillance. One common characteristic of the data acquired in these fields is their uncertain nature. The reasons for uncertainty in data are diverse and ubiquitous shown as follows [1, 2].

1. The uncertainty may be a result of the inherent limitations of the underlying equipment. For example, in sensor and RFID data, uncertainty is due to measurement errors and noises. The output of sensor networks is uncertain because of the noise in sensor inputs and errors in wireless transmission [3].

2. In some applications, such as privacy-preserving data mining [4], it is a requirement that the sensitive data be less precise. For instance, the data is perturbed purposely to preserve the sensitivity of attribute values so that the data can be published [5].

3. In many cases, uncertainty results from the effect of statistical methods such as forecasting and imputation or domain-specific properties and techniques. In the area of probabilistic robotics [6], the behavior or location of a robot is usually
estimated using the Markov or Bayes theory which forms a source of uncertainty due to the limited confidence in the output of predictive models. In information extraction, uncertainty comes from the imperfection of extraction tools and the inherent ambiguity in unstructured text [7].

In some cases, it is possible to eliminate the uncertainties completely. However, this is usually very costly, like manual removal of ambiguous matches in data cleaning. In many cases, complete removal is not even possible. Uncertain data has no place in traditional, precise database management systems like inventory and employee. This has created a need for uncertain data management.

Following this trend, the database and data mining community has investigated extensively the problem of modeling and querying uncertain data [8, 9] in recent decades. Hundreds of research work has been done on this topic [1, 10] and several probabilistic database management systems have been proposed [11–13].

The general approach to manage uncertain data is with a probabilistic model [14], which represents uncertainties using probabilities. According to different granularities, Wang et al. [10] classify uncertainty by three levels: table-based, tuple-based, and attribute-based, with the latter two ones have been widely studied in the literature. The tuple-level uncertainty describes the probability that a tuple exists in the database. On the other hand, the attribute-level uncertainty indicates the probability that a specific value is randomly selected to represent an individual attribute of a tuple from many alternative values. For example, in a database of patients, the disease of a patient can be a cold with a probability of 0.7 and a flu with a probability of 0.3. In this case, the attribute disease is represented by a discrete probability distribution \{0.7, 0.3\}. If the attribute has a continuous domain like the location of a mobile robot, a continuous probability density function such as Gaussian distribution is often utilized to represent this kind of uncertainty.

In this thesis, we focus on the continuous attribute uncertainty and the case where uncertainty is represented by Gaussian distribution. In other words, we assume that uncertain objects stored in the database are represented by Gaussian distributions. Under this setting, we consider query processing over probabilistic data with Gaussian distributions.
Chapter Introduction

We focus our effort on Gaussian distribution because it is one of the most typical probability distributions, and is widely used in statistics, pattern recognition, and machine learning [15, 16]. Especially, in the area of spatio-temporal databases, it is frequently used to represent uncertain location information [14, 17–19].

In Fig. 1.1 and Fig. 1.2 we show examples of one-dimensional and two-dimensional Gaussian distribution, respectively. Generally speaking, it describes the probability that a random point distributes in the space. For example, from Fig. 1.1 in the one-dimensional space, the probability of a point being around the center 0 is about 0.4. The probability decreases as the point spreads from the center. This is the similar in the two-dimensional space in Fig. 1.2. We project the probability surface to a plane and show the decreasing trend with gradient colors. When using Gaussian distribution to represent uncertain location information, it means that the object has the highest probability to be located in the reported location (i.e., the center) and the farther it is, the smaller the probability to be located there becomes.

In the first work, we consider range query, one of the most important queries over spatio-temporal databases. A range query retrieves all the objects that are within the given search range. A range query over uncertain objects, called a probabilistic range query [19, 20], searches for objects within the given search range with probabilities no less than a specified probability threshold. The query object can be either a certain point or an uncertain object represented by a Gaussian distribution. In location-based services, when representing the locations of landmarks by probability distributions like Gaussian distributions,
this kind of query can be used to find surrounding landmarks for a self-navigated mobile robot when building the environment map.

Our second work studies nearest neighbor query or nearest neighbor search, which is also a common type of query in spatio-temporal databases. This query returns the nearest objects to a query point. When extending traditional certain objects to uncertain objects, there are two variations for the new query. The first one, called probabilistic nearest neighbor query [21], utilizes a probability threshold to define qualifying objects in a similar way as the range query. The second one, called expected nearest neighbor search, identifies objects based on their expected distances [22, 23] to the query point. We focus on the second one and consider k-expected nearest neighbor search over uncertain objects represented by Gaussian distributions. An application example is to find potential customers nearby for shops or restaurants.

The third work explores similarity search over Gaussian distributions. Given a database of Gaussian distributions and a query Gaussian distribution, we search the database for top similar data Gaussian distributions. We utilize the Kullback-Leibler divergence (KL-divergence) to measure the similarity between two Gaussian distributions. As in [24], we can represent feature vectors generated in pattern recognition and machine learning using Gaussian distributions and conduct similarity search over them to make interesting and useful findings.

1.2 Research Objectives and Contributions

We have conducted three work on probabilistic data with Gaussian distributions. Our first and second work are motivated by the uncertainty in location information in the area of spatial databases. We represent uncertain locations of objects in the database using Gaussian distributions. In the first work, we assume two kinds of query objects: a certain point and an uncertain location. If the query object is uncertain, we also represent it using a Gaussian distribution. We call them point and Gaussian query object, respectively. Probabilistic range queries by the two kinds of query objects are different and should be handled separately. We show their examples in Fig. 1.3. Here, we define the search
Figure 1.3: Examples of probabilistic range queries

region as the circular area with the query object \( q \) as the center and radius \( \delta \). Given a query object \( q \), a range \( \delta \), and a probability threshold \( \theta \), we search for the objects located within \( \delta \) from \( q \) with probabilities at least \( \theta \).

Although there have been solutions to probabilistic range queries that can handle probabilistic data with Gaussian distributions [25, 26], they are based on specific assumptions and cannot be used to solve the problem in this work. The naïve approach, which sequentially verifies all the objects in the database by computing their exact probabilities, is prohibitively expensive for most real-world applications. This challenge motivates us to develop efficient algorithms to support query processing. We propose a set of filtering techniques to avoid unnecessary computations and a novel indexing method to accelerate query processing. The proposed filtering techniques are based on properties of Gaussian distribution and very effective in reducing computation cost. Due to these techniques, only a small part of objects need to be checked. We develop the indexing method by extending the existing R-tree and improve it based on our analysis of Gaussian distribution. The indexing method organizes all the objects in a structural way so that we can process them by groups efficiently rather than individually.

In the second work, we assume only the point query object. The expected nearest neighbor search returns objects that are close to the query point based on their expected distances. Since most users are interested in the top result objects, we consider \( k \)-expected nearest neighbor search and returns the top-\( k \) results to users. There is no previous work on \( k \)-expected nearest neighbor search over objects represented by Gaussian distributions. Moreover, the naïve approach of performing sequential scan is also computationally expensive for the problem in this work. To explore an efficient solution, we analyze
properties of expected distance on Gaussian distribution mathematically and derive the lower bound and upper bound of this distance. We develop effective filtering techniques using the lower bound and upper bound. The proposed filtering techniques can prune unpromising objects whose lower bound distances are larger than upper bound or expected distances of candidate objects without computing their actual expected distances. To further improve the performance, we utilize R-tree to index all the objects and their lower bound distances and upper bound distances. We propose three novel algorithms to support efficient query processing. The proposed approaches can effectively reduce the number of expensive distance computation.

The third work takes a different view and focuses on the property of Gaussian distribution of being a probability distribution. We use Gaussian distribution as the query object and search for top-$k$ similar Gaussian distributions in the database. The similarity between two Gaussian distributions here is measured by Kullback-Leibler divergence (KL-divergence) [27], which is a representative measure for quantifying the similarity between two probability distributions. Since KL-divergence is a non-metric measure which violates the properties of a standard distance function in metric spaces, existing solutions for metric spaces cannot be employed to solve this problem. Although there have been several studies on searching in non-metric spaces [28–31], their problem settings are different from ours. The naïve solution of sequential scan is again not suitable for this problem. To this end, We propose two types of approaches utilizing the notions of rank aggregation [32] and skyline queries [33]. The first type presorts all objects in the database on their attributes and computes result objects by merging candidates from each presorted list. The second one transforms the problem to the computation of dynamic skyline queries. We extend and modify the branch-and-bound skyline (BBS) algorithm, which is proposed to answer skyline queries, and develop a novel algorithm to solve this problem. We propose novel algorithms to support efficient query processing by modifying and extending existing algorithms.

Our contributions are summarized as follows:

- We formalize two types of probabilistic range queries with respect to the point and Gaussian query object.
• For the two types of probabilistic range queries, we propose effective filtering techniques and a novel indexing method to support efficient query processing.

• We formally define the problem of $k$-expected nearest neighbor search over objects represented by Gaussian distributions.

• We analyze mathematically properties of the expected distance on Gaussian distribution and derive the lower bound and upper bound of this distance.

• We propose three novel approaches to improve the efficiency of $k$-expected nearest neighbor search.

• We formalize the problem of top-$k$ similarity search based on KL-divergence over Gaussian distributions, and analyze mathematically the properties of KL-divergence between two Gaussian distributions.

• We propose two types of approaches to improve the efficiency of query processing using the notion of rank aggregation and skyline queries.

• We demonstrate the efficiency and effectiveness of our approaches through comprehensive experimental performance studies.

1.3 Related Work

The field of uncertain data management has received considerable attention from many researchers since its growing up in recent decades. Managing uncertainty poses a number of unique challenges on several fronts. The two broad issues are those of modeling the uncertain data, and then leveraging it to work with various applications [1]. In [34, 35], several issues and working models for uncertain data have been discussed. The other issue is that of adapting data management and mining applications for uncertain data. There have been a number of excellent surveys and books discussing uncertain data management [1, 9, 10, 34, 36, 37]. Li et al. [38] provides a cross-disciplinary view of uncertainty processing activities by different communities.
Several models have been proposed to incorporate uncertain objects in databases [14], including the fuzzy model [39], the evidence-oriented model [40], and the probabilistic model. Moreover, probabilistic graphical models [41] are proposed to represent complex dependencies among uncertain data. These models are mainly different in the semantics and complexities of the data in the underlying applications [35]. In the area of database, the general approach to manage uncertain data is the use of probabilistic model, which represents uncertainties using probabilities. According to different granularities of uncertainty, the probabilistic model can be further classified into three categories: table-based, tuple-based, and attribute-based [10, 14], with the latter two ones have been widely studied in the literature.

Specifically, a table-based approach concerns the “coverage” of a table, i.e., the percentage of tuples present in a table [42]. On the other hand, a tuple-based solution associates each tuple with a probability indicating the likelihood that the tuple exists in the table [43]. An attribute-based method represents the attribute of a tuple using a probability distribution [3, 19, 44, 45]. The probability distribution can be either a discrete one, which describes a set of possible values with occurring probabilities [3, 44], or a continuous one such as a Gaussian distribution [19, 45]. The former case is often dealt with using the possible worlds semantics [46], while the latter one depends on properties of specific probability density functions. Our work falls into the latter category.

There have been diverse studies on adapting data management and mining activities for uncertain data. Several probabilistic database management systems have been proposed, including Trio [11], MayBMS [12], and Orion [13]. In the area of spatio-temporal databases, a large amount of work has been done to support efficient processing of various queries such as probabilistic range queries [14, 19, 45], probabilistic nearest neighbor queries [21], probabilistic join queries [47], and probabilistic skyline queries [48]. Their continuous versions have also been investigated for moving objects to either query historical records, i.e., trajectories [49, 50], or monitor current activities [51, 52]. In addition, there are other versions incorporated with query types like threshold queries [53], top-k queries [54], ranking queries [55], aggregate queries [56], and reverse queries [57], and versions in different environments such as distributed domains [58], indoor space [59], constrained space [60], and road networks [61]. We show the diagram of our research...
work in this thesis in Fig. 1.4. Our first work in Chapter 2 combines probabilistic range queries with the threshold query predicate, and our second work in Chapter 3 integrates probabilistic nearest neighbor search with top-k queries. On the other hand, our third work in Chapter 4 considers both similarity search and top-k queries.

Uncertain data management has also been studied in other fields such as data integration [62], data streams [63], XML data [64], graph data [65], sensor networks [66] and biological images [67]. Moreover, mining uncertain data has received much attention from multiple research communities including database, data mining, and information retrieval. A number of algorithms for clustering uncertain data [68, 69], outlier detection [70] and frequent pattern mining [71] from uncertain data are proposed.

### 1.4 Thesis Organization

The rest of the thesis is organized as follows. In Chapter 2 we introduce the probabilistic range querying over Gaussian objects. Then we present the k-expected nearest neighbor search over Gaussian objects in Chapter 3. Chapter 4 describes the top-k similarity search over Gaussian distributions based on KL-divergence. Finally, in Chapter 5 we conclude our studies and present future directions.
Chapter 2

Probabilistic Range Querying over Gaussian Objects

2.1 Introduction

In the field of uncertain data management, probabilistic range query is an important problem for processing uncertain data in real-world applications such as mobile robotics, location-based services, and sensor networks. A probabilistic range query returns all the data objects that appear within the given search region with probabilities no less than a given probability threshold.

For instance, consider a self-navigated mobile robot moving in a wireless environment. The robot builds a map of the environment by observing nearby landmarks via devices such as sonars and laser range finders. Due to the inherent limitation brought about by sensor accuracy and signal noises, the location information acquired from measuring devices is not always precise. At the same time, the robot also conducts probabilistic localization \[6\] to estimate its own location autonomously by integrating its movement history and the landmark information. This may cause impreciseness in the location of the robot, too. In consequence, probabilistic queries have evolved to tackle such impreciseness as “find landmarks that are located within 5 meters from my current location with probabilities at least 80%”. 

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Typically for such applications, uncertain objects are stored in the database and associated with probability distributions. Multi-dimensional *Gaussian distribution* is one of the commonly used probability distributions for such a purpose. It is widely adopted in statistics, pattern recognition [16], and localization in robotics [6].

In this work we study the case where the locations of data objects are uncertain, whereas the location of the query object is either exact or uncertain. Specifically, data objects are described by Gaussian distributions with different parameters to indicate their respective uncertainty. A query object can be either a certain point in the multi-dimensional space or an uncertain location represented by a multi-dimensional Gaussian distribution. We solve the probabilistic range query problem according to the above setup.

There have been solutions to probabilistic range queries that can handle uncertain data represented by Gaussian distributions, yet based on specific assumptions. For example, U-tree [25] assumes that each uncertain object exists within a *pre-defined uncertainty region*. It constructs an index structure for all the objects based on this region to reduce the number of candidates that require expensive numerical integration. However, for some application scenarios it is not easy to decide a suitable extent of the uncertainty region for a real-world object. Gauss-tree [26] assumes that the Gaussian distribution must be *independent* in each dimension. When these assumptions are violated, the solutions no longer work. In this work, we solve these problems with generic Gaussian distributions without any of these assumptions, i.e., an object can locate in an infinite space as opposed to U-tree, and have correlations between dimensions as opposed to Gauss-tree.

A straightforward approach to this problem is to compute the *appearance probability* [14] for each data object and output it if this probability is no less than the threshold. However, the probability computation usually requires costly numerical integration for the accurate result [25], rendering it prohibitively expensive to compute for all the data objects and check if the query constraint is satisfied. Thus, such computations should be reduced as much as possible.

To this end, we propose filtering techniques to generate a set of candidate data objects and compute integrations only for these candidates. Equipped with the filtering techniques, an R-tree-based indexing method is proposed to accelerate query processing. The index
structure is inspired by the idea of TPR-tree [72], in which the minimum bounding boxes (MBBs) vary with time. The difference is that in our index a parent MBB not only varies with the probability threshold but also tightly encloses all the child MBBs.

In our preliminary work [19], we proposed query processing algorithms for probabilistic range queries, assuming that only the location of the query object is uncertain and represented by a Gaussian distribution, but data objects are certain multi-dimensional points. An R-tree can be used to manage these certain data points to process queries, which is different from the situation here. In this work, we extend the uncertainty to data objects and propose novel solutions. We develop several effective filtering techniques by analyzing properties of Gaussian distribution thoroughly and design a non-trivial indexing method that can handle multiple online queries.

A precedent report of this work has appeared in [73]. The approach proposed in [73] approximates the Gaussian distribution with an upper-bounding function. An R-tree-like hierarchical index structure was proposed and an exponential summary function was defined to cover multiple upper-bounding functions. Nevertheless, the summary function is so sensitive to the child functions that it will become drastically large if the bounded Gaussian distributions are sparsely distributed in the space or one of them has large variances, leading to loose bounding in the index structure and weak filtering power. This work is an extension of our previous work [45]. We have extended the algorithms and conducted additional experiments.

Our contributions are summarized as follows:

1. We formalize two types of probabilistic range queries with respect to the query object: a certain point and an uncertain location represented by a Gaussian distribution, while data objects are represented by Gaussian distributions with different parameters.

2. For the two types of queries, we propose several effective filtering techniques to identify promising data objects and prune unpromising ones.

3. We design a novel R-tree-based index structure to support probabilistic range queries on Gaussian objects.
4. We demonstrate the efficiency of our approach through a comprehensive experimental performance study using real datasets.

The rest of this chapter is organized as follows. Section 2.2 defines our problem. We present our filtering strategies in Section 2.3. Section 2.4 describes our index structure. Experimental results and analyses are covered by Section 2.5. Section 2.6 reviews related work. Section 2.7 concludes this chapter.

2.2 Problem Definition

In this section, we first define Gaussian objects, and then define probabilistic range queries using two types of query objects: certain point objects and uncertain Gaussian objects.

2.2.1 Gaussian Objects

The Gaussian distribution, also known as normal distribution, is a continuous probability distribution defined by a bell-shaped probability density function in the one-dimensional case. In this work, we assume that data objects are modeled by Gaussian distributions in a $d$-dimensional space. A point $x$ is in a $d$-dimensional numerical space, namely, $x = (x^1, \ldots, x^d)^T$.

**Definition 2.1** (Gaussian objects). A Gaussian object $o$ is represented by its possible locations (points) and the probability density it appears at each location. Formally, the probability density that $o$ is located at $x_o$ is captured by a $d$-dimensional Gaussian probability density function

$$p_o(x_o) = \frac{1}{(2\pi)^{d/2}|\Sigma_o|^{1/2}} \exp \left[ -\frac{1}{2}(x_o - \mu_o)^T \Sigma_o^{-1} (x_o - \mu_o) \right]. \quad (2.1)$$

$\mu_o$ is the mean location (center) of $o$. $\Sigma_o$ is a $d \times d$ covariance matrix. $|\Sigma_o|$ and $\Sigma_o^{-1}$ are the determinant and the inverse of $\Sigma_o$, respectively. $x^T$ denotes the transposition of the vector $x$. 
2.2.2 Probabilistic Range Queries on Gaussian Objects

Given a database of Gaussian objects $D$, a query object $q$, a distance threshold $\delta$, and a probability threshold $\theta$, a probabilistic range query (PRQ) on Gaussian objects retrieves all the data objects $o \in D$ such that the distance between $o$ and $q$ is no more than $\delta$ with probabilities no less than $\theta$. In this work, we consider two types of query objects for $q$:

1. The query object is a point object, namely,
   \[ q = (x_{q1}, x_{q2}, \ldots, x_{qd})^t. \]

2. The query object is a Gaussian object, namely,
   \[ p_q(x_q) = \frac{1}{(2\pi)^{d/2} |\Sigma_q|^{1/2}} \exp \left[ -\frac{1}{2} (x_q - \mu_q)^t \Sigma_q^{-1} (x_q - \mu_q) \right]. \]

The probabilistic range query with a point query object (PRQ-P) is formally defined as

\[ \text{PRQ-P}(D, q, \delta, \theta) = \{ o \mid o \in D, \Pr(\|x_o - q\| \leq \delta) \geq \theta \}, \]

where $\|x_o - q\|$ represents the Euclidean distance between $x_o$ and $q$. We call the region consisting of the points with distances no more than $\delta$ from the query object the query region, $QR$ for short. $\Pr(\|x_o - q\| \leq \delta)$, the probability that $o$ lies within $QR$, is computed by

\[ \Pr(\|x_o - q\| \leq \delta) = \int \chi_\delta(x_o, q) \cdot p_o(x_o) dx_o, \quad (2.2) \]

where

\[ \chi_\delta(x_o, q) = \begin{cases} 1, & \|x_o - q\| \leq \delta; \\ 0, & \text{otherwise}. \end{cases} \quad (2.3) \]

The integration in Eq. (2.2) is not in a closed-form and hence cannot be computed directly. Numerical solutions such as Monte Carlo methods can be employed to evaluate the probability. We use the importance sampling [74] in the interest of efficiency. Specifically, we generate $x_o$ as per the probability function $p_o(x_o)$, and increment the count when
Figure 2.1: PRQ-P query

Eq. (2.3) is satisfied. Finally, the probability can be obtained by dividing the count by the number of samples generated. Generally speaking, however, Monte Carlo methods are accurate only if the number of samples is sufficiently large (at the order of $10^6$) [25]. Therefore, the probability computation induces expensive runtime cost.

Figure 2.1 illustrates a PRQ-P query in a 2-dimensional space. The Gaussian object $o$ exists in the space with decreasing probability densities as it spreads from the center $\mu_o$. A PRQ-P query finds the Gaussian objects located in the proximity of the query point with a required probability. Computing the probability using Eq. (2.2) corresponds to integrating the probability density function of $o$ within the shaded area around $q$.

Similar to PRQ-P, the probabilistic range query with a Gaussian query object (PRQ-G) is defined as

$$
\text{PRQ-G}(D, q, \delta, \theta) = \{o \mid o \in D, \Pr(\|x_o - x_q\| \leq \delta) \geq \theta\},
$$

where $\Pr(\|x_o - x_q\| \leq \delta)$ is computed by

$$
\Pr(\|x_o - x_q\| \leq \delta) = \int\int \chi_\delta(x_o, x_q) \cdot p_o(x_o) \cdot p_q(x_q) d{x_o} d{x_q},
$$

(2.4)

where

$$
\chi_\delta(x_o, x_q) = \begin{cases} 
1, & \|x_o - x_q\| \leq \delta; \\
0, & \text{otherwise.}
\end{cases}
$$
To compute the integration in Eq. (2.4), although we can simply generate random numbers for the two Gaussian distributions \( p_o(x_o) \) and \( p_q(x_q) \), respectively, a more efficient method is shown in [73]. It constructs a 2\(d\)-dimensional Gaussian distribution by combining the two \(d\)-dimensional Gaussian distributions. Then the probability can be computed by running the Monte Carlo method.

### 2.3 Filtering Based on Approximated Region

A naïve algorithm to answer PRQ-P or PRQ-G queries is to pair the query object with each data object and perform integration check with either Eq. (2.2) or Eq. (2.4). However, due to the overhead of the integration, the algorithm becomes prohibitively expensive for large datasets. So we develop our approach based on the filter-and-refine paradigm, i.e., to obtain a set of candidate objects and then compute the integration only for the candidates.

In this section, we first introduce the notion of \( \rho \)-region that leverages the two thresholds \( \delta \) and \( \theta \), and then propose the \( \rho \)-region-based filtering techniques to handle PRQ-P and PRQ-G queries.

#### 2.3.1 \( \rho \)-Region

**Definition 2.2 (\( \rho \)-region).** Consider a Gaussian object \( o \) and the integration of its probability density function \( p_o(x_o) \) over an ellipsoidal region \( (x_o - \mu_o)^T \Sigma_o^{-1} (x_o - \mu_o) \leq r^2 \). Let \( r_o \) be the value of \( r \) within which the integration equals \( \rho \):

\[
\int_{(x_o - \mu_o)^T \Sigma_o^{-1} (x_o - \mu_o) \leq r^2} p_o(x_o) dx_o = \rho.
\]

We call the ellipsoidal region

\[
(x_o - \mu_o)^T \Sigma_o^{-1} (x_o - \mu_o) \leq r_o^2
\]

the \( \rho \)-region of \( o \).
In Fig. 2.1, the area encompassed by the dotted ellipsoidal curve shows a $\rho$-region. Because the probability density of a Gaussian distribution decreases as we move away from the center of the object, if the query object is distant enough from the center, its probability within $QR$ will not reach $\theta$. In other words, it is possible to determine whether a data object can satisfy the query condition by deriving a suitable $\rho$-region with $\theta$ (the method is introduced in Section 2.3.3 and 2.3.4) and examining whether its $\rho$-region intersects $QR$.

To compute $r_\rho$ with a given $\rho$, we borrow the approach proposed in our previous work [19]. It transforms the integration over an ellipsoidal region to an integration over a $d$-dimensional spherical region. By assigning $\mu_o = 0$ and $\Sigma_o = I$ in Eq. (2.1), we have the normalized Gaussian distribution

$$p_{\text{norm}}(x) = N(0, I) = \frac{1}{(2\pi)^{d/2}} \exp \left[ -\frac{1}{2} ||x||^2 \right].$$

Based on this probability density function, the following property can be derived.

**Property 1.** [19] Consider integration of $p_{\text{norm}}(x)$ over $||x||^2 \leq r^2$. For a given $\rho$ ($0 < \rho < 1$), let $\tilde{r}_\rho$ be the radius within which the integration becomes $\rho$:

$$\int_{||x||^2 \leq \tilde{r}_\rho^2} p_{\text{norm}}(x) dx = \rho.$$ 

Then $r_\rho = \tilde{r}_\rho$ holds.

The preceding property indicates that we can compute $\tilde{r}_\rho$ and hence $r_\rho$ for a given $\rho$ value. Therefore, we can construct a $(\rho, r_\rho)$-table offline (numerical integration is necessary) and obtain the $\rho$-region by looking up the corresponding $r_\rho$ from this table. If there is no matched entry for a given $\rho$, we conservatively return the corresponding $r_\rho$ of the smallest value greater than $\rho$, so the correctness of the result can be guaranteed.

The ellipsoidal shape of a $\rho$-region renders it difficult to quickly examine whether the $\rho$-region intersects $QR$ as well as develop an indexing scheme based on prevalent spatial indexes such as R-tree. Hence we will study the minimum bounding box (MBB) which tightly bounds the $\rho$-region.
2.3.2 Minimum Bounding Box of $\rho$-Region

Figure 2.2 shows the MBB of a $\rho$-region of a 2-dimensional Gaussian object $o$. Let $w_j$ denote the length of its edge along the $j$-th dimension. The following property holds [19].

Property 2. The value of $w_j$ ($j = 1, \ldots, d$) is given as

$$w_j = \sigma_j r_{\rho},$$

where $\sigma_j$ corresponds to the standard deviation in the $j$-th dimension

$$\sigma_j = \sqrt{(\Sigma_o)_{jj}},$$

where $(\Sigma_o)_{jj}$ represents the $(j, j)$-th element of $\Sigma_o$.

For a data object $o$, since $\sigma_j$ can be calculated from the covariance matrix $\Sigma_o$, the scale of the MBB is determined uniquely by $r_{\rho}$, and hence $\rho$. Consequently, in order to establish the filtering conditions utilizing the MBBs, it is essential to explore the relation between $\rho$ and the probability threshold $\theta$. Next we will present our filtering techniques for PRQ-P and PRQ-G, respectively.

2.3.3 Filtering Policies for PRQ-P Queries

Our filtering policies to process PRQ-P queries are divided into two cases: $\theta < 0.5$ and $\theta \geq 0.5$. 
Case 1 ($\theta < 0.5$): Consider the four data objects $o_1$, $o_2$, $o_3$, $o_4$ shown in Fig. 2.3(a). $bb_i(\rho)$ denotes the MBB of $o_i$’s $\rho$-region. First, let’s consider $o_4$. Since the probability that $o_4$ is located inside its $\rho$-region is $\rho$, the probability of being outside the $\rho$-region’s MBB, is definitely less than $1 - \rho$. Furthermore, given the line symmetry of the Gaussian distribution, the probability that $o_4$ exists inside $QR$ is at most $(1 - \rho)/2$. Hence, if $\rho = 1 - 2\theta$, and $bb_4(\rho)$ and $QR$ do not overlap, the probability that $o_4$ lies within $QR$ must be less than $\theta$. Then, for objects $o_1$, $o_2$ and $o_3$, we check and find their MBBs intersect $QR$, so they become candidates. In summary, when $\theta < 0.5$, a data object is a candidate only if its $bb_i(1 - 2\theta)$ intersects $QR$.

Case 2 ($\theta \geq 0.5$): We show our idea in Fig. 2.3(b). For the probability that a data object exists inside $QR$ to reach $\theta$, its mean location should lie within $QR$. Thus, $o_2$ and $o_4$ can be pruned, whereas $o_1$ and $o_3$ are considered as candidates.

Moreover, for all candidates, let $\rho' = \theta$ and compute their $bb_i(\rho')$’s. If $QR$ fully contains a $bb_i(\rho')$ (e.g., $o_3$), the probability that this object lies within $QR$ is definitely greater than $\theta$. We validate it as a result without computing the numerical integration.

Summarizing the two cases, the filtering condition for PRQ-P is: (1) $\theta < 0.5$, $bb_i(\rho)$ ($\rho = 1 - 2\theta$) intersects $QR$ for pruning, and $bb_i(\rho')$ ($\rho' = \theta$) is contained by $QR$ for validation. (2) $\theta \geq 0.5$, $\|\mu - q\| < \delta$ for pruning, and $bb_i(\rho')$ ($\rho' = \theta$) is contained by $QR$ for validation.

2.3.4 Filtering Policies for PRQ-G Queries

For PRQ-G queries, since both the query object $q$ and the data object $o_i$ are Gaussian distributions, we obtain both of their MBBs of $\rho$-regions. We also consider two cases for filtering: $\theta < 0.75$ and $\theta \geq 0.75$.

Case 1 ($\theta < 0.75$): As shown in Fig. 2.3(c) assume $q$ and $o_i$ are independent in the space. When the minimum distance between the two MBBs is exactly $\delta$, the maximal probability of $\|x_i - x_q\| \leq \delta$ is given by the following lemma (the proof is in A.1):

Lemma 2.3. If $\text{MinDist}(bb_i(\rho), bb_q(\rho)) \geq \delta$, $Pr(\|x_i - x_q\| \leq \delta) < (3 - 2\rho - \rho^2)/4$. 

Let \( (3 - 2\rho - \rho^2)/4 = \theta \), i.e., \( \rho = 2 \sqrt{1 - \theta} - 1 \). \( o_i \) can be excluded from the candidate set if the minimum distance between \( bb_i(\rho) \) and \( bb_q(\rho) \) is no less than \( \delta \).

**Lemma 2.4.** If \( \|\mu_o - \mu_q\| \geq \delta \), \( \Pr(\|x_o - x_q\| \leq \delta) < 0.75 \).

**Case 2** (\( \theta \geq 0.75 \)): Based on Lemma 2.4 (the proof is in A.2), an object can be pruned if the distance between its center and that of the query object is no less than \( \delta \).

Moreover, let \( \rho' = \sqrt{\theta} \), if the maximum distance of \( bb_i(\rho') \) and \( bb_q(\rho') \) is less than \( \delta \), as shown in Fig. 2.3(d), the probability of \( \|x_o - x_q\| \leq \delta \) is greater than \( \rho^2 \), i.e., \( \theta \). In this case, we validate it as a result without computing the exact probability by numerical integration.

The filtering condition for PRQ-G is summarized as: (1) \( \theta < 0.75 \), \( \text{MinDist}(bb_i(\rho), bb_q(\rho)) < \delta \) \( (\rho = 2 \sqrt{1 - \theta} - 1) \) for pruning, and \( \text{MaxDist}(bb_i(\rho'), bb_q(\rho')) < \delta \) \( (\rho' = \sqrt{\theta}) \) for validation. (2) \( \theta \geq 0.75 \), \( \|\mu_o - \mu_q\| < \delta \) for pruning, and \( \text{MaxDist}(bb_i(\rho'), bb_q(\rho')) < \delta \) \( (\rho' = \sqrt{\theta}) \) for validation.
Chapter 2
Probabilistic Range Querying over Gaussian Objects

2.4 Indexing Data Objects

2.4.1 The Index Structure

The filtering conditions introduced in the previous section need to know the value of \( \theta \) and hence \( \rho \) to generate candidates. In order not to scan all the data objects and compute the MBBs of the \( \rho \)-regions on the fly with the given \( \theta \), an immediate solution is to index the MBBs for a sufficiently large \( \rho_{\text{max}} \). Because the MBB with a larger \( \rho \) always subsumes the one with a smaller \( \rho \), it can support all the queries if the \( \rho \) values computed from \( \theta \) satisfy the condition \( \rho \leq \rho_{\text{max}} \). However, the efficiency of the index is compromised for small \( \rho \) values. This method serves as a baseline algorithm (we use an R-tree to index MBBs and name it FR-tree), and will be compared in the experiment with the indexing technique we are going to present.

Inspired by the TPR-tree \[72\], we propose an R-tree-based index structure which stores the MBBs in a parametric fashion. It works for arbitrary probability thresholds and range thresholds, and there is no need to assume the two thresholds are given prior to index construction. The MBBs can be dynamically computed as we traverse the index. Furthermore, the MBB of a node (at both leaf and non-leaf levels) tightly encloses all its children’s MBBs regardless of the \( \theta \) value, as opposed to the TPR-tree within which all child MBBs are bounded in a loose manner.

Our index is a balanced, multi-way tree organized in the structure of an R-tree. Each entry in a leaf node contains a data object in the form of \( o_i = (id_i, \mu_i, \Sigma_i) \), where \( id_i \) is the data object id, \( \mu_i \) and \( \Sigma_i \) are the mean location and the covariance matrix of the Gaussian object, respectively. In a non-leaf node, each entry has a pointer to a child node and an MBB enclosing all the MBBs within the child node.

Consider an object \( o_i \) with mean location \( (x'_1, \ldots, x'_d) \). Its MBB is a bounding box parameterized with \( r_p \) (MBB is denoted as the exactly computed bounding box at a specific \( r_p \) hereafter). From Eq. (2.5), the extent of the bounding box (BB) in the \( j \)-th dimension can be represented by

\[
[x'_j - w'_j, x'_j + w'_j] = [x'_j - \sigma'_j r_p, x'_j + \sigma'_j r_p]. \tag{2.6}
\]
Figure 2.4: BB with varying $\rho$

Seeing the BBs grow with $r_\rho$, in order to tightly bound the BBs of child nodes at every $r_\rho$, it is necessary to search each dimension for the leftmost and the rightmost BBs under varying $\rho$. We illustrate this problem in Fig. 2.4(a). It shows the changing BB that encloses the BBs of four 2-dimensional objects’ $\rho$-regions as $r_\rho$ increases. When $r_\rho$ is less than $r_1$, the left edge is determined by $o_1$, and it becomes $o_3$ when $r_\rho$ exceeds $r_1$. The right bound is determined by $o_4$ when $r_\rho < r_2$, and $o_2$ otherwise.

Figure 2.4(b) shows how the four BBs change horizontally with $r_\rho$. For each object, a pair of symmetrical lines describe the left and the right coordinations of its BB. The lines have different slopes due to the difference in their standard deviations. The bold polylines illustrate the left and right coordinations of the outer enclosing BB. Hence, the problem becomes how to find the bold polylines. To this end, a BB can be represented by several segments with respect to $r_\rho$.

We store in the index the $j$-th dimension of a BB in the form of $((x'_j, \sigma'_j, r_1), \ldots, (x'_k, \sigma'_k, +\infty))$. For example, for the four objects in Fig. 2.4(a), the left and the right coordinations of the BB are $((x'_1, \sigma'_1, r_1), (x'_4, \sigma'_4, +\infty))$ and $((x'_4, \sigma'_4, r_2), (x'_2, \sigma'_2, +\infty))$, respectively.

We can find all the segments in the $j$-th dimension through a sort on the coordinations first and then a linear scan from the object whose standard deviation has a larger value in the $j$-th dimension. The time complexity is $O(n \log n)$, where $n$ is the number of its child nodes. The number of segments in a BB is at most $n$.

---

1 Our experiments show the average number of segments in a BB is 2–3. In case of many segments, we allow users to specify a query probability threshold range $[\theta_{\text{min}}, \theta_{\text{max}}]$ to reduce the number of segments and hence the index size.
We note the difference between our index and TPR-tree: (1) The bounding boxes of TPR-tree change towards one direction in a rate (velocity), while our bounding boxes change towards two opposite directions symmetrically with $r_p$. (2) The bounding boxes of TPR-tree are tight only if an update occurs, while our bounding boxes are always tight.

### 2.4.2 Query Processing with the Index

Algorithm 1 and Algorithm 2 show the procedures of query processing of PRQ-P (the algorithms for PRQ-G are similar and thus omitted due to the space limitation). To process a query, $\theta$ is converted to $\rho$ and $\rho'$, and then $r_\rho$ and $r_{\rho'}$ with the pre-computed $(\rho, r_\rho)$ table (line 1–2). A first-in-first-out queue $Q$ is employed to maintain the nodes we are going to expand. Starting with the root node, we compare $QR$ (MBB of $q$ for PRQ-G) with the node MBB, and check the filtering condition. To compute a node MBB, given $r_\rho$, we scan the stored $j$th-dimension of its BB and find $\alpha$ such that $r_{\alpha-1} \leq r_\rho < r_\alpha$. Then the extent of its MBB in the $j$-th dimension can be computed through Eq. (2.6) using $x_\alpha$, $\sigma_\alpha$, and $r_\rho$.

At the non-leaf level (line 7–18), if a node MBB at $r_{\rho'}$ satisfies the validation condition, i.e., it is contained by $QR$ (for PRQ-G, the maximum distance of two MBBs is less than $\delta$), we retrieve all the objects within this node and push them into the result set directly; otherwise, we probe each child entry. If $\theta < 0.5$ (0.75 for PRQ-G), we check whether each child MBB at $r_\rho$ intersects $QR$ (for PRQ-G, whether the minimum distance between each child MBB and the MBB of $q$ is less than $\delta$). On the other hand, when $\theta \geq 0.5$ (0.75 for PRQ-G), we obtain the MBB which bounds leftmost and rightmost means in each dimension, and use this more compact MBB as the new child MBB for filtering checking for both PRQ-P and PRQ-G.

At the leaf level (line 19–30), we also examine whether a node MBB at $r_{\rho'}$ satisfies the validation condition first, so as to push its child entries into the result set as early as possible and reduce the processing cost. If not, each child entry of this node is processed as discussed in Section 2.3.3 (for PRQ-G, Section 2.3.4). When $\theta < 0.5$ (0.75 for PRQ-G), for each child entry, if its MBB at $r_\rho$ intersects $QR$ (for PRQ-G, the minimum distance between each child MBB and the MBB of $q$ is less than $\delta$), we identify this data object
using Algorithm 2. We check whether the validation condition is satisfied. The qualified data objects are inserted directly into the result set. Other data objects are regarded as candidates for further verification through integration. When \( \theta \geq 0.5 \) (0.75 for PRQ-G), for each child entry, if its mean location is inside \( QR \), it is checked by Algorithm 2 in the same way.

Figure 2.5 shows a query processing example for PRQ-P with \( \theta = 0.3 \). \( r_{0.4} \) and \( r_{0.3} \) are obtained from the \((\rho, r_{\rho})\) table. MBBs of all nodes in Fig. 2.5 are at \( r_{0.4} \). At first, the root \( R \) is popped from \( Q \). Since the MBB of \( R \) cannot satisfy the validation condition, we have to check its child nodes \( n_1, n_2 \) and \( n_3 \). Only \( n_1 \) and \( n_2 \) are pushed into \( Q \) because the MBB of \( n_3 \) does not intersect \( QR \). Then we continue with the next level. For \( n_1 \), its MBB at \( r_{0.4} \) is not contained by \( QR \), so its child entries cannot be all validated as results. Since its MBB at \( r_{0.3} \) intersects \( QR \), we probe into its child entries and find that \( o_1 \) can be pruned. Then \( o_2 \) is validated as a result object and \( o_3 \) becomes a candidate further processed through integration. For \( n_2 \), since its MBB at \( r_{0.3} \) is completely within \( QR \), its entries \( o_4 \) and \( o_5 \) are inserted into the result set directly without referring to their MBBs.

2.5 Experiments

2.5.1 Experimental Setup

Three real datasets are used in our experiments. MG and LB are two 2-dimensional datasets of Montgomery and Long Beach road networks (39K and 52K, respectively).\(^2\)

\(^2\)http://www.census.gov/geo/www/tiger
Algorithm 1 PRQ-P\((q, \delta, \theta)\)

1: \(\rho \leftarrow 1 - 2\theta, \rho' \leftarrow \theta\)
2: \(r_\rho \leftarrow TableLookup(\rho), r_{\rho'} \leftarrow TableLookup(\rho')\)
3: \(QR \leftarrow QueryRegion(q, \delta), ResultSet = \emptyset, CandidateSet = \emptyset\)
4: \(Q \leftarrow \text{Root}, N \leftarrow 0\)
5: \(\text{while } Q \neq \emptyset \text{ do}\)
6: \(N \leftarrow Q.pop()\)
7: \(\text{if } N \text{ is an internal node then} \quad \triangleright \text{at the non-leaf level}\)
8: \(\quad \text{if } bb_N(r_\rho) \text{ IsContainedBy } QR \text{ then}\)
9: \(\quad \quad \text{Insert all the objects within } N \text{ into } ResultSet\)
10: \(\quad \text{else}\)
11: \(\quad \quad \text{if } \theta < 0.5 \text{ then}\)
12: \(\quad \quad \quad \text{for each child } N_i \text{ in } N \text{ do}\)
13: \(\quad \quad \quad \quad \text{if } bb_N(r_\rho) \text{ Intersects } QR \text{ then}\)
14: \(\quad \quad \quad \quad \quad \text{Push } N_i \text{ into } Q\)
15: \(\quad \quad \text{else} \quad \triangleright \theta \geq 0.5\)
16: \(\quad \quad \text{for each child } N_i \text{ in } N \text{ do}\)
17: \(\quad \quad \quad \text{if } meanbb_N \text{ Intersects } QR \text{ then}\)
18: \(\quad \quad \quad \quad \text{Push } N_i \text{ into } Q\)
19: \(\quad \quad \text{else} \quad \triangleright \text{at the leaf level}\)
20: \(\quad \quad \text{if } bb_N(r_{\rho'}) \text{ IsContainedBy } QR \text{ then}\)
21: \(\quad \quad \quad \text{Insert all the objects within } N_i \text{ into } ResultSet\)
22: \(\quad \quad \text{else}\)
23: \(\quad \quad \quad \text{if } \theta < 0.5 \text{ then}\)
24: \(\quad \quad \quad \quad \text{for each child } N_i \text{ in } N \text{ do}\)
25: \(\quad \quad \quad \quad \quad \text{if } bb_i(r_\rho) \text{ Intersects } QR \text{ then}\)
26: \(\quad \quad \quad \quad \quad \quad \text{IdentifyObjects}(N_i)\)
27: \(\quad \quad \quad \text{else} \quad \triangleright \theta \geq 0.5\)
28: \(\quad \quad \quad \text{for each child } N_i \text{ in } N \text{ do}\)
29: \(\quad \quad \quad \quad \quad \text{if } mean(N_i) \text{ IsInside } QR \text{ then}\)
30: \(\quad \quad \quad \quad \quad \quad \text{IdentifyObjects}(N_i)\)

Airport is a 3-dimensional dataset containing latitudes, longitudes and elevations of 41K airports in the world\(^3\). All datasets are normalized to \([0, 1000]\)^d. LB is used by default.

We generate PRQ-P and PRQ-G queries randomly. The probability threshold \(\theta\) lies within \([0.01, 0.99]\), and the query range \(\delta\) is chosen from \([10, 100]\) for MG and LB, and \([100, 200]\) for Airport randomly. We also generate covariance matrices for both data Gaussian objects and query Gaussian objects randomly.

\(^3\)http://www.ourairports.com/data
We design two baseline approaches for experimental evaluation. Both of them and our proposed approach return the same exact result. We check their correctness before each comparison. One baseline approach is to sequentially scan the dataset and compute integrations required for obtaining result probabilities. We name it Scan and evaluate our filtering techniques by comparing query processing time and candidate number with it. The other baseline approach indexes the MBBs of the $\rho$-region with $\rho_{\text{max}} = 0.99$. Because the MBB with a larger $\rho$ always subsumes the one with a smaller $\rho$, it can support all the queries if the $\rho$ values computed from $\theta$ satisfy $\rho \leq \rho_{\text{max}}$. We equip this index with our basic filtering techniques and name it FR-tree (Filtering + R-tree), and evaluate our index structure by comparing filtering time and I/O access with it. Our index is referred to as G-tree.

We implement the index structure by extending the spatial index library SaiL\[^4\][75]. It is a generic framework that integrates spatial and spatio-temporal index structures and supports user-defined datatypes and customizable spatial queries. We conduct experiments using a PC with Intel Core 2 Duo CPU E8500 (3.16GHz), RAM 4GB, running Fedora 12. We construct an index of all data objects for both PRQ-P and PRQ-G, and store it in the secondary memory.

### 2.5.2 Query Performance Evaluation

The average query response time of 200 queries on LB is shown in Table 2.1. It can be seen that the query response time of Scan is 769 times that of G-tree for PRQ-P (293 for PRQ-G). Another observation is that the time spent on probability integration is almost equal to the overall response time. This indicates that the integration dominates the

[^4]: http://libspatialindex.github.com/
Chapter 2. Probabilistic Range Querying over Gaussian Objects

<table>
<thead>
<tr>
<th>Query / Algorithm</th>
<th>Overall</th>
<th>Integration</th>
</tr>
</thead>
<tbody>
<tr>
<td>PRQ-P / G-tree</td>
<td>0.157</td>
<td>0.154</td>
</tr>
<tr>
<td>PRQ-P / Scan</td>
<td>120.764</td>
<td>120.692</td>
</tr>
<tr>
<td>PRQ-G / G-tree</td>
<td>0.809</td>
<td>0.806</td>
</tr>
<tr>
<td>PRQ-G / Scan</td>
<td>236.725</td>
<td>236.577</td>
</tr>
</tbody>
</table>

**Table 2.1: Query response time on LB (seconds)**

![Bar charts showing performance metrics](image)

**Figure 2.6: Filtering and indexing performance**

Overall query processing and is computationally expensive. Consequently, it is important to reduce candidates which need to perform the integration as much as possible.

Among 50,747 objects in LB, the average candidate number of G-tree is 97 for PRQ-P (269 for PRQ-G). The number of validated objects by integration is 65 for PRQ-P (156 for PRQ-G). So for PRQ-P 67% (58% for PRQ-G) of the candidates identified by our approach are real results. This demonstrates the effectiveness of our proposed filtering techniques. In the sequel, we exclude the integration part from query processing and focus on evaluating the filtering and indexing performance of FR-tree and G-tree.
We run the two algorithms to process 10K queries on the three datasets and show the average filtering time and I/O access of PRQ-P (resp. PRQ-G) in Fig. 2.6(a)–2.6(c) (resp. Fig. 2.6(b)–2.6(d)). For PRQ-P, the average filtering time of G-tree is 61.6% of that of FR-tree on the three datasets, because the average I/O access of G-tree is 92.2% less than that of FR-tree, though the segmented bounding boxes in G-tree are more complex to process than those in FR-tree. The reduction on PRQ-G is more substantial. The average filtering time of G-tree on MG, LB and Airport is 45% less than that of FR-tree. The I/O access of G-tree of three datasets is 6.5% that of FR-tree.

As a $\rho_{\text{max}}$ is adopted to process queries with any $\theta$, the bounding boxes in FR-tree are very loose. This causes more I/O accesses and increases filtering time. In contrast, since the bounding boxes in G-tree are constructed in a parametric fashion, they can be calculated dynamically for arbitrary $\theta$ and hence are very compact. Another interesting observation is that the I/O access almost resembles the candidate number, indicating most I/Os are spent on retrieving objects.

Figure 2.6(e)–2.6(f) show the candidate ratio of PRQ-P and PRQ-G, which is calculated by dividing the candidate number by the total number of objects. The candidate number of FR-tree and G-tree is the same since we equip FR-tree with our filtering techniques. The candidate ratio is around 2‰ for PRQ-P and 5‰ for PRQ-G on the three datasets. This reveals that only a very small percentage of data objects will become candidates owing to our filtering techniques.

**Varying Dataset Size.** To evaluate the scalability of our approach, we extract 20%, 40%, 60%, 80% and 100% of LB dataset randomly and show the filtering time and I/O access of two methods in Fig. 2.7(a)–2.7(b) on PRQ-P queries. The performance on PRQ-G queries reveals a similar trend and hence is omitted due to the space limitation. As the dataset size becomes larger, the filtering time and I/O access of the two methods almost increase linearly. G-tree displays a steady increasing trend and always outperforms FR-tree.

As shown in Fig. 2.7(c)–2.7(d) in spite of the varying dataset size $|\mathcal{D}|$, the candidate ratio of PRQ-P retains 2‰ and 5.2‰ for PRQ-G, demonstrating the steadiness and scalability of our approach with respect to the dataset size.
Varying Query Range. We vary the query range $\delta$ from 10 to 100 and show the performance on PRQ-P queries in Fig. 2.8(a)–2.8(b). The performance on PRQ-G queries is similar and hence omitted. As $\delta$ increases, FR-tree consumes much more time and I/O accesses on filtering. In contrast, G-tree exhibits much slower increasing trends. Figure 2.8(c)–2.8(d) show that the candidate ratio of both PRQ-P and PRQ-G also increases with $\delta$, but for PRQ-P it is only 3.7‰ (9.4‰ for PRQ-G) even if $\delta$ achieves 100.

Varying Probability Threshold. We vary $\theta$ from 0.1 to 0.9 and show the performance in
Fig. 2.9(a)–2.9(f) for both PRQ-P and PRQ-G queries. For PRQ-P, the filtering time and I/O access of both FR-tree and G-tree decrease gradually with \( \theta \) when \( \theta < 0.5 \) (0.75 for PRQ-G). When \( \theta \) exceeds 0.5 (0.75 for PRQ-G), the filtering time slightly rebounds. This is consistent with our filtering conditions as discussed in Section 2.3.3 and Section 2.3.4.

When \( \theta < 0.5 \) (0.75 for PRQ-G), \( \rho = 1 - 2\theta \) for PRQ-P (for PRQ-G, \( \rho = 2\sqrt{1-\theta} - 1 \) if \( \theta < 0.75 \)), so \( \rho \) decreases when \( \theta \) moves towards larger values, and bounding boxes shrink. As a result, most of non-candidates can be filtered quickly and less I/O accesses are needed, and hence it accelerates filtering.

On the contrary, when \( \theta \geq 0.5 \) (0.75 for PRQ-G), the pruning condition becomes \( ||\mu_o - q|| < \delta \) for PRQ-P (for PRQ-G, \( ||\mu_o - \mu_q|| < \delta \)). So all the figures have turn points at 0.5 for PRQ-P (0.75 for PRQ-G). At the same time, bounding boxes computed for validation which assigns \( \rho' = \theta \) for PRQ-P (for PRQ-G, \( \rho' = \sqrt{\theta} \)) enlarge as \( \theta \) increases. So more and more nodes and objects will become candidates, leading to the slightly rising of the filtering time and I/O access. The reason also accounts for the trend of G-tree on candidate ratio in Fig. 2.9(e)–2.9(f).
Despite the variation of $\theta$, G-tree constantly outperforms FR-tree. In the case of PRQ-P, the filtering time of FR-tree amounts to 1.8 times that of G-tree on average and 15.8 times on average for I/O access. This contrast is more evident on PRQ-G, where the filtering time of FR-tree is 2 times that of G-tree on average and 20.1 times on average for I/O access.

**Varying Dimensionality.** We also study the impact of dimensionality $d$ using randomly generated synthetic datasets with the size 20K and the query range within [100, 200]. Figure 2.10(a) – 2.10(d) show the scalability of FR-tree and G-tree against $d$ for PRQ-P. The figures of filtering time and I/O access for PRQ-G have similar trends and are omitted. As shown in Fig. 2.10(a), the filtering time of FR-tree reduces constantly with increasing $d$ because the object density decreases with $d$. This can be confirmed by the decreasing trend of I/O access of PRQ-P in Fig. 2.10(b) and the candidate ratio of both PRQ-P and PRQ-G in Fig. 2.10(c) – 2.10(d).

It is also observed that the filtering performance of FR-tree begins to exceed that of G-tree at $d = 5$. The explanation is that, candidates become fewer as object density decreases, and hence the operation of comparing the query region with node bounding boxes dominates the filtering procedure. While FR-tree’s MBBs can be obtained directly from the R-tree, G-tree needs to compute the MBBs with the segments in the index. As $d$ increases, the decreasing object density can lower the processing cost. However, the increasing $d$ can also induce more effort in query processing since the bounding box computation needs to be done for more dimensions. The above two factors result in the fluctuating trend of G-tree’s filtering time in Fig. 2.10(a).

**Index construction.** We evaluate the index construction on the Airport dataset. The node capacities of the indexes are selected to optimize query performance for both FR-tree and G-tree. The index size of FR-tree is 6.94MB, and the construction time is 4.3 seconds on average. G-tree has a size of 6.14MB, slightly smaller than FR-tree due to the different entry style of an object from FR-tree. At the leaf level of FR-tree, for each object, besides the bounding box at $p_{max}$, the covariance matrix has to be stored additionally in order to carry out our filtering techniques. In contrast, in G-tree the bounding box of an object is formed by its mean and standard deviation in each dimension. In addition to the
bounding box, only the covariances between dimensions rather than the whole covariance matrix need to be stored. G-tree takes 59.7 seconds on average to build. Although G-tree needs more construction time, considering the superior query performance and the index construction can be done offline, the index construction is in an affordable manner.

2.6 Related Work

**Uncertain Data Management.** We focus on research work in the area of uncertain data management that is closely related to our work. A number of approaches for managing uncertain data have been proposed. Early research primarily focused on queries in a moving object database model [76–79]. The solutions to several types of probabilistic queries were proposed in [44], including probabilistic range queries, where their target is merely the one-dimensional case.

A range query processing method for the case where both data objects and query object are imprecise was proposed in [80]. But they assume that each object exists within a rectangular area. Zheng et al. [81] modeled a fuzzy object by a fuzzy set where each element is characterized by its probability of membership (the sum of all probabilities is not necessarily one). For efficient query processing, they proposed the notion of $\alpha$-cut,
the subset of elements whose probabilities are no less than a user-specified probability threshold $\alpha$, to filter elements of the fuzzy object. Although we also exploit the idea of filtering region ($\rho$-region), their rationale of computing the $\alpha$-cut is different from ours.

Gauss-tree [26] was proposed as an index structure for Gaussian distributions. It assumes all Gaussian distributions are probabilistically independent in each dimension. This imposes heavy restriction on the generality of the approach and the overall accuracy of the query result is limited. In [82], Lian et al. proposed a generic framework to tackle the local correlations among uncertain data.

**Indexing Uncertain Data for Range Queries.** Agarwal et al. [17] presented various indexing structures on uncertain data that support range queries in the one-dimensional case. Tao et al. proposed U-tree [25] to process probabilistic range queries in a multi-dimensional space, where uncertain objects are assumed to follow arbitrary probability distributions within uncertainty regions. Zhang et al. proposed a quadtree-based index called U-Quadtree [83] for range searching on multi-dimensional uncertain data. They mainly focused on representing uncertainty by discrete instances inside a minimum bounding box. The difference from our work is that we take advantage of specific properties of Gaussian distribution and index uncertain objects distributed in an infinite space.

**Spatial Data Indexing.** The traditional spatial database has been well studied and many indexing methods have been proposed [84–86] to support spatial query processing. R-tree [85] and its extension R*-tree [84], indexing objects by deriving their minimum bounding rectangles (MBRs), are two of the well-known ones. TPR-tree [72] and TPR*-tree [87] were proposed to index moving objects. However, none of these indexes can be applied directly on the Gaussian objects to support the queries studied in this work.

### 2.7 Summary

In this work, we studied probabilistic range queries over uncertain data. We assumed that the location of the query object is either fixed or follows a multi-dimensional Gaussian distribution. The locations of data objects are represented by Gaussian distributions.
Given these assumptions, we defined two types of probabilistic range queries with respect to the query object. We proposed effective filtering techniques to reduce costly exact probability computation as much as possible. Furthermore, we proposed a novel R-tree-based index structure to expedite query processing. Extensive experiments on real datasets demonstrate the efficiency and effectiveness of our proposed approach.
Chapter 3

$k$-Expected Nearest Neighbor Search over Gaussian Objects

3.1 Introduction

In recent years, the advances in computing devices and technologies have been calling for the attention of researchers to develop novel solutions for emerging new problems. For instance, the location information obtained from mobile devices such as a mobile phone is usually uncertain due to privacy protection, delays in location update from users, noise, etc. In the area of database, there has been a large amount of work on representing uncertain location information by probabilistic models and proposing efficient solutions for query processing [10]. In particular, Gaussian distribution is frequently used to represent this kind of probabilistic location information since it is one of the most typical probability distributions, and is widely used in statistics, pattern recognition, and machine learning [15, 16]. For example, our first work in Chapter 2 presents probabilistic range queries over objects represented by Gaussian distributions.

On the other hand, as one of the commonest queries over location information, the distance-based nearest neighbor search has applications in numerous fields such as databases, pattern recognition, cluster analysis, and recommendation systems. Given a query point
Figure 3.1: An example

$q$, this query searches for closest objects to $q$ in a set of objects. There have been considerable efforts made to extend nearest neighbor search over traditional location information to probabilistic location information. One representative example is the *expected distance* [22, 23], which defines the distance over probabilistic location information. Following this trend, in this work, we represent probabilistic location information by Gaussian distributions and assume that the closeness between each Gaussian object and the query point is measured by their *expected squared Euclidean distance* [23], which is frequently used in many areas such as pattern mining and cluster analysis. Under this setting, we consider the problem of *$k$-expected nearest neighbor search over Gaussian objects*.

Although there has been a considerable amount of research work on searching over Gaussian distributions [20, 26], to the best of our knowledge, we are the first to consider the problem of *$k$-expected nearest neighbor search over objects represented by Gaussian distributions*. Given a database $D$ of objects represented by Gaussian distributions, a query point $q$, and a constant $k$, we search $D$ for the top-$k$ objects having smallest distances with $q$. As shown in Fig. 3.1, an application example is to find nearby mobile users to a given query location such as a shop or a restaurant. Since the location of a mobile user is uncertain, it is a common practice to represent uncertain locations using Gaussian distributions in the area of spatial databases [20]. The restaurant or shop may consider sending coupons or notifications to a number of nearest users for promotion.

The expected distance used in this work is defined using an integral and is normally computed by numerical integration such as the Monte Carlo methods. This kind of methods requires a sufficiently large number of samples (e.g., $10^5$) to ensure accuracy. Hence,
it will lead to very high time cost if we process queries by comparing distances of all objects one by one in real time. This naïve solution is intolerable for the vast majority of real-world applications which demand immediate responses. What is more, the world of today is being flooded with streams of large data and is in the age of big data. This calls for novel approaches that can handle a large dataset and support efficient query processing.

To find an efficient solution, we analyze properties of expected distance on Gaussian distribution mathematically and derive the lower bound and upper bound of this distance. Meanwhile, we employ the filter-and-refine paradigm to accelerate query processing. By filtering, we prune unpromising objects whose lower bound distances are larger than upper bound or expected distances of candidate objects without computing their actual distances. In refinement, we compute exact distances of candidate objects and finally return the top-$k$ smallest ones. To further improve the performance, we utilize R-tree to index objects and their lower bound distances and upper bound distances. We propose three novel algorithms to support efficient query processing. The experimental result demonstrates that our proposed approaches can achieve great efficiency and are applicable to real-world applications.

We summarize our contributions as follows.

1. We formally define the problem of $k$-expected nearest neighbor search over objects represented by Gaussian distributions.

2. We analyze mathematically properties of the expected distance on Gaussian distribution and derive the lower bound and the upper bound of this distance.

3. We propose three novel approaches to improve the efficiency of query processing.

4. We demonstrate the efficiency of our approaches through a comprehensive experimental performance study.

The rest of this chapter is organized as follows. We formally define the problem in Section 3.2. Then in Section 3.3, we analyze expected distance on Gaussian distribution and derive its lower bound and upper bound. We propose three approaches in Section 3.4.
Experimental results and analyses are presented in Section 3.5. We review related work in Section 3.6. Finally, Section 3.7 concludes this chapter.

### 3.2 Problem Definition

In this work, we consider the problem of $k$-nearest neighbor search over Gaussian objects based on expected distance. We assume that all objects stored in the database are represented by Gaussian distributions with different parameters, i.e., their average vectors and covariance matrices are not the same. The query objects are fixed points in the same space.

A Gaussian distribution in the multi-dimensional space is represented by

$$p(x) = \frac{1}{(2\pi)^{d/2}|\Sigma|^{1/2}} \exp \left[-\frac{1}{2}(x - \mu)^T \Sigma^{-1} (x - \mu)\right]$$

where $\mu$ denotes the average vector and $\Sigma$ denotes the covariance matrix. $|\Sigma|$ and $\Sigma^{-1}$ represent the determinant matrix and reverse matrix of $\Sigma$, respectively. $(x - \mu)^T$ represents the transposition of $x - \mu$. For simplicity, we call objects represented by Gaussian distribution Gaussian objects afterwards.

Here, we focus on the squared Euclidean distance (SQED) since it is frequently used in many areas such as pattern mining and cluster analysis. The expected distance between a Gaussian object $o$ and a query point $q$ based on SQED (called ESQED) is

$$ESQED(o, q) = \int ||x - q||^2 \cdot p(x)dx.$$ (3.1)

We formally define the problem as follows.

**Definition 3.1.** Given a database of Gaussian objects $D = \{o_1, \ldots, o_n\}$, a query point $q$, a constant $k$, the problem of $k$-Expected Nearest Neighbor Search over Gaussian objects searches $D$ for $k$ Gaussian objects that are nearest to $q$ based on Eq. (3.1).
The distance in Eq. (3.1) cannot be computed directly and is normally evaluated by numerical integration using the Monte Carlo methods. We employ the importance sampling [88] for efficiency. Specifically, we generate a sample $x$ based on $p(x)$ and sum its SQED with $q$. The ESQED is the summed SQED divided by the number of samples generated. This computation requires a sufficiently large number of samples (e.g., $10^5$) to ensure accuracy. Therefore, it will induce very high computation cost.

A straightforward solution is to do a sequential scan over the database and compute their distances with each query point. This method is obviously too time-consuming for real-world applications. In the preliminary experiments, we generate 100,000 two-dimensional Gaussian objects and use 100,000 samples for numerical integration. The average runtime of 100 queries is 1623.4 seconds, which is rather expensive and intolerable for the most of users.

In the following sections, we utilize the filter-and-refine paradigm to accelerate query processing. Specifically, we filter objects that are unlikely to become the result without computing their ESQEDs and refine candidate objects to obtain the final result. The filtering is to prune objects whose lower bound distances are larger than upper bound or expected distances of candidate objects. By refinement, we compute the ESQED of each candidate object and choose the top-$k$ objects with the smallest ESQED as the result. Hence, it is important to derive the lower bound and the upper bound of ESQED (Section 3.3) and develop effective filtering algorithms (Section 3.4).

### 3.3 Lower bound and Upper bound

In [23], the following lemma is proved.

**Lemma 3.2.** Given an object $p$ with $f(x)$ as its probability density function and $\overline{p}$ as its centroid, for any point $q$ we have

$$
\int ||p - q||^2 \cdot f(x)dx = ||\overline{p} - q||^2 + \int ||x - \overline{p}||^2 \cdot f(x)dx.
$$
Based on this lemma, we can obtain

$$ES\ QED(o, q) = ||\mu - q||^2 + \int ||x - \mu||^2 \cdot p(x) dx.$$  

In the subsequent discussion, we denote the second part as $\Delta^2$, i.e.,

$$\Delta^2 = \int ||x - \mu||^2 \cdot p(x) dx,$$

and derive its lower bound and upper bound. According to [19], the lower bound and upper bound of Gaussian distribution is as follows:

$$p^{\perp}(x) = \frac{1}{(2\pi)^{d/2}|\Sigma|^{1/2}} \exp \left[ -\frac{||x||^2}{2\lambda^{\perp}} \right]$$

$$p^{\top}(x) = \frac{1}{(2\pi)^{d/2}|\Sigma|^{1/2}} \exp \left[ -\frac{||x||^2}{2\lambda^{\top}} \right]$$

where $\lambda^{\perp}$ and $\lambda^{\top}$ are the minimum and maximum eigenvalues of $\Sigma$, respectively. Thus, the lower bound of $\Delta^2$, denoted as $\Delta^2_{\text{min}}$, is

$$\Delta^2 \geq \int ||x - \mu||^2 \cdot \frac{1}{(2\pi)^{d/2}|\Sigma|^{1/2}} \exp \left[ -\frac{||x - \mu||^2}{2\lambda^{\perp}} \right] dx$$

$$= \frac{d \cdot (\lambda^{\perp})^{1+d/2}}{|\Sigma|^{1/2}} = \Delta^2_{\text{min}},$$

and its upper bound $\Delta^2_{\text{max}}$ is

$$\Delta^2 \leq \int ||x - \mu||^2 \cdot \frac{1}{(2\pi)^{d/2}|\Sigma|^{1/2}} \exp \left[ -\frac{||x - \mu||^2}{2\lambda^{\top}} \right] dx$$

$$= \frac{d \cdot (\lambda^{\top})^{1+d/2}}{|\Sigma|^{1/2}} = \Delta^2_{\text{max}}$$

For the proof, please refer to Appendix B.1.

Correspondingly, we obtain the lower bound (LB) and upper bound (UB) of ESQED:

$$LB(o, q) = ||\mu - q||^2 + \Delta^2_{\text{min}} \leq ES\ QED(o, q) \leq ||\mu - q||^2 + \Delta^2_{\text{max}} = UB(o, q).$$  \hspace{1cm} (3.2)
3.4 Proposed Approaches

In this section, we propose three approaches for efficient query processing using the filter-and-refine paradigm described in Section 3.2. However, it still seems cumbersome and inefficient to compare the lower bound and upper bound distance of each Gaussian object. Our idea is to employ an R-tree to index all Gaussian objects so that most unpromising objects can be filtered by groups instead of individually. Moreover, we observe that $\Delta^2_{\min}$ and $\Delta^2_{\max}$ involve costly matrix decomposition but they do not depend on queries. Based on this observation, we consider precomputing $\Delta^2_{\min}$ and $\Delta^2_{\max}$ offline to speed up online query processing.

In the first approach, we first precompute $\Delta^2_{\min}$ and $\Delta^2_{\max}$ for each Gaussian object and then insert each $d$-dimensional Gaussian object into an R-tree as a $(d+1)$-dimensional region $[(\mu, \Delta^2_{\min}), (\mu, \Delta^2_{\max})]$. Here, $(\mu, \Delta^2_{\min})$ is the lower left coordinate and $(\mu, \Delta^2_{\max})$ is the upper right coordinate of the region. In this way, we can easily compute the lower bound and upper bound distance of each Gaussian object using Eq. (3.2) when queries come.

At the same time, we associate each Gaussian object with its covariance matrix so that we can compute its ESQED directly if it becomes a candidate. It should be noted that we only need to store the lower triangular part or the upper triangular part of each covariance matrix since it is symmetric. In other words, for a $d \times d$ covariance matrix, we store $d(d+1)/2$ out of its $d^2$ elements.

For an R-tree node $N$ with a region of $[(\mu_{\min}, \Delta^2_{\min}), (\mu_{\max}, \Delta^2_{\max})]$, its lower bound and upper bound distance can be computed similarly using Eq. (3.2):

$$LB(N, q) = \|\mu - q\|^2_{\min} + \Delta^2_{\min} \leq ESQED(N, q) \leq \|\mu - q\|^2_{\max} + \Delta^2_{\max} = UB(N, q). \quad (3.3)$$

Below we use an example dataset of six two-dimensional Gaussian objects in Table 3.1 to describe our algorithms. $\mu_{i,1}$ (resp. $\mu_{i,2}$) denotes the average of each object $o_i$ in the first (resp. second) dimension. $\lambda_{i,1}$ (resp. $\lambda_{i,2}$) denotes the eigenvalue of the variance matrix of each object $o_i$ in the first (resp. second) dimension. $\Delta^2_{i,\min}$ (resp. $\Delta^2_{i,\max}$) denotes $\Delta^2_{\min}$ (resp. $\Delta^2_{\max}$) of each object $o_i$. 
Table 3.1: An example dataset

<table>
<thead>
<tr>
<th>$o_1$</th>
<th>$\mu_{i,1}$</th>
<th>$\mu_{i,2}$</th>
<th>$\lambda_{i,1}$</th>
<th>$\lambda_{i,2}$</th>
<th>$\Delta_{i,\min}^2$</th>
<th>$\Delta_{i,\max}^2$</th>
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</thead>
<tbody>
<tr>
<td>$o_1$</td>
<td>0.5</td>
<td>4.0</td>
<td>0.5</td>
<td>1.0</td>
<td>0.71</td>
<td>2.83</td>
</tr>
<tr>
<td>$o_2$</td>
<td>2.9</td>
<td>5.8</td>
<td>1.0</td>
<td>1.0</td>
<td>2.00</td>
<td>2.00</td>
</tr>
<tr>
<td>$o_3$</td>
<td>4.0</td>
<td>6.0</td>
<td>1.5</td>
<td>0.5</td>
<td>0.58</td>
<td>5.20</td>
</tr>
<tr>
<td>$o_4$</td>
<td>6.7</td>
<td>2.0</td>
<td>1.0</td>
<td>1.5</td>
<td>1.63</td>
<td>1.84</td>
</tr>
<tr>
<td>$o_5$</td>
<td>8.5</td>
<td>3.0</td>
<td>0.5</td>
<td>0.5</td>
<td>1.00</td>
<td>1.00</td>
</tr>
<tr>
<td>$o_6$</td>
<td>9.0</td>
<td>1.5</td>
<td>0.5</td>
<td>1.0</td>
<td>0.71</td>
<td>2.83</td>
</tr>
</tbody>
</table>

In Fig. 3.2, we show the image and structure of the constructed R-tree on the example dataset and an example query. Each black solid point represents the average value in the first dimension $\mu_{i,1}$ and in the second dimension $\mu_{i,2}$ of each object $o_i$ in the $(\mu_{i,1}, \mu_{i,2})$ plane. The example query is the black hollow point $q = (5.0, 5.0)$ with $k = 2$. We can easily compute $LB(N_1, q) = 0 + 1^2 + 0.58 = 1.58$, $LB(N_2, q) = 7.60$, $LB(o_1, q) = 21.96$, and $UB(o_5, q) = 7.20$. Notice that $UB(o_2, q) = 7.05$ because $\lambda_{2,1} = \lambda_{2,2}$. It means $ES\ QED(o_2, q) = 7.05$. In other words, the ESQED of a Gaussian object can be computed without numerical integration if its matrix covariance has the same eigenvalues in all dimensions. The same is to $o_5$, $UB(o_5, q) = LB(o_5, q) = 17.25$, i.e., $ES\ QED(o_5, q) = 17.25$.

Algorithm 3 shows the algorithm of our first approach. Since it is based on precomputing, $LB$, and $UB$, we call it PLUB afterwards. Entries (R-tree nodes or Gaussian objects) are processed in the ascending order of their lower bound distances. We utilize a priority queue $Q$ to maintain candidate entries. After expanding the root node of the R-tree, entries in $Q$ are $\{(1.58, N_1), (7.60, N_2)\}$. Then the algorithm continues to expand $N_1$. Since
Algorithm 3 The PLUB algorithm

1: Input: $R$-tree, $q$, $k$ \hspace{1cm} \triangleright \text{Initialize the priority queue } Q, \text{ the candidate set } C, \text{ and the result set } R$
2: $Q \leftarrow \text{Root of } R$-tree, $C \leftarrow \emptyset$, $R \leftarrow \emptyset$;
3: \textbf{while } $Q$ is not empty \textbf{do}
4: \hspace{1cm} $N \leftarrow Q$.top(); $Q$.pop();
5: \hspace{1cm} \textbf{if } $N$ is a leaf node \textbf{then}
6: \hspace{2cm} \textbf{for } each Gaussian object $o_i$ in $N$ \textbf{do}
7: \hspace{3cm} \textbf{if } $C$ has less than $k$ candidates \textbf{then}
8: \hspace{4cm} Insert $o_i$ with $LB(o_i, q)$ and $UB(o_i, q)$ into $C$;
9: \hspace{3cm} \textbf{else} \hspace{0.5cm} \triangleright C has $k$ or more than $k$ candidates
10: \hspace{4cm} \textbf{if } $LB(o_i, q)$ is smaller than the $k$-th smallest UB in $C$ \textbf{then}
11: \hspace{5cm} Insert $o_i$ with $LB(o_i, q)$ and $UB(o_i, q)$ into $C$;
12: \hspace{3cm} \textbf{else} \hspace{0.5cm} \triangleright N$ is an index node
13: \hspace{4cm} \textbf{if } $C$ has less than $k$ candidates \textbf{then}
14: \hspace{5cm} \textbf{for } each child entry $N_i$ in $N$ \textbf{do}
15: \hspace{6cm} Insert $N_i$ with $LB(o_i, q)$ into $Q$;
16: \hspace{4cm} \textbf{else} \hspace{0.5cm} \triangleright C has $k$ or more than $k$ candidates
17: \hspace{5cm} \textbf{for } each child entry $N_i$ in $N$ \textbf{do}
18: \hspace{6cm} \textbf{if } $LB(N_i, q)$ is smaller than the $k$-th smallest UB in $C$ \textbf{then}
19: \hspace{7cm} Insert $N_i$ with $LB(o_i, q)$ into $Q$;
20: \hspace{4cm} \textbf{if } $C$ has $k$ or more than $k$ candidates and $LB(Q$.top$(q)$ is not smaller than the $k$-th smallest UB in $C$ \textbf{then}
21: \hspace{5cm} \text{break;}$
22: \text{Compute the ESQEDs of the top-$k$ candidates with the smallest LB in $C$ and insert them with their ESQEDs into $R$}
23: \textbf{for } each of the rest candidates $o_j$ in $C$ \textbf{do}
24: \hspace{1cm} \textbf{if } $LB(o_j, q)$ is smaller than the largest ESQED in $R$ \textbf{then}
25: \hspace{2cm} Compute $o_j$’s ESQED and replace it with $o_i$ and $o_j$’s ESQED;
26: \textbf{return } R;$

$N_1$ is a leaf node and the candidate set $C$ is empty, we insert $o_1$, $o_2$, and $o_3$ and their LBs and UBs into $C$. We obtain $C = \{(2.58, 7.20, o_3), (7.05, 7.05, o_2), (21.96, 24.08, o_1)\}$ and $Q = \{(7.60, N_2)\}$. The next top entry of $Q$, $N_2$, has an LB that is larger than the second largest UB in $C$, i.e., $7.60 > 7.20$, which is the second smallest upper bound distance in $C$. Thus, we finish processing $Q$ and continue to update the result set $R$ using $C$.

We first obtain the ESQED of $o_3$ by numerical integration since it LB is the smallest. For $o_2$, we do not have to compute its ESQED by numerical integration because we can directly obtain $ESQED(o_2, q) = 7.05$ as described above. By this time, $R = \{(3.99, o_3), (7.05, o_2)\}$. Since the LB of the last object $o_1$ in $C$, 21.96, is larger than 7.05,
we return \( R \) and terminate the algorithm.

The second approach, called PLB, only utilizes the lower bound. After precomputing \( \Lambda^2_{\text{min}} \) for each Gaussian object, we insert each \( d \)-dimensional Gaussian object into an R-tree as a \((d + 1)\)-dimensional point \((\mu, \Lambda^2_{\text{min}})\). We can easily compute the lower bound distance of each Gaussian object using Eq. (3.2) when queries come. We also associate each Gaussian object with its covariance matrix so that we can compute its ESQED directly if it becomes a candidate. The lower bound distance of an R-tree node can be computed in the same way as PLUB using Eq. (3.3).

In the PLB algorithm, instead of inserting each candidate Gaussian object \( o_i \) with \( LB(o_i, q) \) and \( UB(o_i, q) \) into \( C \) (Line 8), we immediately compute its ESQED and insert \( o_i \) with \( ESQED(o_i, q) \) into \( C \). Moreover, we compare the LB of \( o_i, N_i \), and \( Q.top() \) with the \( k \)-th smallest ESQED rather than the \( k \)-th smallest UB in \( C \) (Line 10, 18, and 20). In Line 11, we replace the candidate having the \( k \)-th smallest ESQED of \( C \) with \( o_i \) if \( ESQED(o_i, q) \) is smaller. Finally, since \( C \) has already had the top-\( k \) candidates with the smallest ESQEDs in this time we skip Line 22–25 and return \( C \) directly in Line 26.

Continuing with the example dataset in Table 3.1 and the example query of \( q = (5.0, 5.0) \) with \( k = 2 \), in the PLB algorithm we expand the root node of the R-tree (the R-tree constructed by PLB is almost the same to that of PLUB in Fig. 3.2 and obtain \( Q = \{(1.58, N_1), (7.60, N_2)\} \). When expanding \( N_1 \), we computes ESQEDs of \( o_1, o_2, \) and \( o_3 \) in turn and update \( C = \{(3.99, o_3), (7.05, o_2)\} \) and \( Q = \{(7.60, N_2)\} \). Then the algorithm will be terminated since the LB of the next top entry in \( Q \) is larger than the second largest ESQED in \( C \), i.e., \( 7.60 > 7.05 \). Finally, we return \( C = \{(3.99, o_3), (7.05, o_2)\} \).

The idea of the PLB algorithm is that, using ESQED rather than UB would help to prune more entries since ESQED is always smaller and thus tighter than UB. At the same time, however, it may induce more ESQED computations since it immediately computes the ESQED of each candidate. For instance, we compute three times of ESQED using PLB while only one time using PLUB in the running example. We will compare the performance of PLUB and PLB in the experiment section.
The third approach does not rely on precomputing and directly utilizes the average of each Gaussian object, and is called AVG. Notice that by Eq. \((3.2)\) we can obtain the least lower bound (LLB) of ESQED as follows:

\[
\text{LLB}(o, q) = \|\mu - q\|^2 < \|\mu - q\|^2 + \Delta_{\text{min}}^2 \leq \text{ESQED}(o, q).
\]  (3.4)

The least lower bound is looser than the lower bound but it allows us to filter unpromising objects using the simple average directly without caring about the complicated covariance matrix. We insert each \(d\)-dimensional Gaussian object into an R-tree as a \(d\)-dimensional point \(\mu\). We compute the least lower bound distance of each Gaussian object using Eq. \((3.4)\) when queries come. We also associate each Gaussian object with its covariance matrix as we do previously. The least lower bound distance of an R-tree node is the minimum least lower bound distance of all its child nodes.

Algorithm 4 shows the AVG algorithm. It is very similar to the PLB algorithm except that it uses LLB as the search key instead of LB. We describe the algorithm by using the previous example in PLUB and PLB. The R-tree constructed by AVG is similar to the one in Fig. 3.2 other than that in this case the R-tree is two-dimensional. We first expand the root node of the R-tree and obtain \(Q = \{(1.0, N_1), (6.89, N_2)\}\). Then we expand \(N_1\) and computes ESQEDs of \(o_1\), \(o_2\), and \(o_3\) in turn. Meanwhile, we update \(C = \{(3.99, o_3), (7.05, o_2)\}\) and \(Q = \{(6.89, N_2)\}\). Next, since the LLB of the next top entry in \(Q\) is smaller than the second largest ESQED in \(C\), i.e., \(6.89 < 7.05\), the algorithm cannot be terminated as PLUB and PLB do. In other words, we have to further expand \(N_2\) and examine \(o_4\), \(o_5\), and \(o_6\) through computing their ESQEDs. Finally, we return \(C = \{(3.99, o_3), (7.05, o_2)\}\).

The AVG algorithm does not have to do any precomputing and the R-tree constructed by it has a lower dimension than of PLUB and PLB. As the performance of R-tree deteriorates with higher dimensions, we think this lower dimensional R-tree of AVG may compensate the cost of more ESQED computations caused by the looser least lower bound. We will verify this idea in the experiment section.
Algorithm 4 The AVG algorithm

1: Input: R-tree, $q$, $k$  
$\triangleright$ Initialize the priority queue $Q$, the candidate set $C$
2: $Q \leftarrow$ Root of R-tree, $C \leftarrow \emptyset$;
3: while $Q$ is not empty do
4:   $N \leftarrow Q$.top(); $Q$ .pop();
5:   if $N$ is a leaf node then
6:     for each Gaussian object $o_i$ in $N$ do
7:        if $C$ has less than $k$ candidates then
8:           Insert $o_i$ with $ESQED(o_i, q)$ into $C$;
9:        else $\triangleright$ $C$ has $k$ or more than $k$ candidates
10:           if $LLB(o_i, q)$ is smaller than the $k$-th smallest $ESQED$ in $C$ then
11:              Replace it with $o_i$ and $ESQED(o_i, q)$;
12:        else $\triangleright$ $N$ is an index node
13:          if $C$ has less than $k$ candidates then
14:             for each child entry $N_i$ in $N$ do
15:                Insert $N_i$ with $LLB(o_i, q)$ into $Q$;
16:          else $\triangleright$ $C$ has $k$ or more than $k$ candidates
17:             for each child entry $N_i$ in $N$ do
18:                if $LLB(N_i, q)$ is smaller than the $k$-th smallest $ESQED$ in $C$ then
19:                   Insert $N_i$ with $LLB(o_i, q)$ into $Q$;
20:          if $C$ has $k$ candidates and $LLB(Q$.top$(o_i, q)$ is not smaller than the $k$-th smallest
21:             ESQED in $C$ then
22:             break;
23: return $C$;

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Testing Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k$</td>
<td>1, 5, 10, 15, 20, 25, 30, 35, 40, 45, 50</td>
</tr>
<tr>
<td>data size</td>
<td>10K, 100K, 1M, 10M, 100M</td>
</tr>
<tr>
<td>dimension</td>
<td>2, 3, 4, 5, 6</td>
</tr>
</tbody>
</table>

Table 3.2: Parameters for testing

3.5 Experiments

3.5.1 Experimental Settings

We generate Gaussian distributions randomly for experiments. Each average value is generated from $(0, 1000)$, and each variance value is generated from $(0, 100)$. We check the effect of $k$, data size, and $d$, i.e., the number of result, the size of dataset, the dimensionality, on the performance of each approach. The parameters tested in our experiments and their default values (in bold) are summarized in Table 3.2.
We compare the performance of our three proposed approaches: AVG, PLB, and PLUB. During preprocessing, we construct R-trees and store them in the secondary memory. In each experiment, we run 100 queries (generated randomly) and use the average runtime and I/O access for performance evaluation. All experiments are conducted using a workstation with Intel Xeon (R) CPU E3-1241 v3 (3.50GHz), RAM 16GB, running Ubuntu 12.10 LTS. Below we analyze the effect of each parameter on the performance of query processing and discuss the merits and demerits of the proposed approaches.

### 3.5.2 Effect of $k$

We show the average runtime and I/O access of the three approaches when varying $k$ in Fig. 3.3(a) and Fig. 3.3(b). Their average runtime and I/O access all increases slowly with a larger $k$. This is because that we need to do more computations to retrieve more objects. As we can see from the two figures, PLUB runs the fastest, but its I/O access is also the highest. The reason is that most of objects accessed by PLUB will be pruned by upper bounds of candidate objects using their lower bounds and we only need to compute ESQEDs for a small part of them. Since the computation of ESQED consumes quite high time cost, this helps PLUB to achieve the best efficiency. In addition, when $k$ is larger than 5, AVG performs better than PLB because the two-dimensional R-tree of AVG is more effective and retrieves less objects than the three-dimensional R-tree of PLB.
3.5.3 Effect of Data Size

As shown in Fig. 3.4(a) and Fig. 3.4(b), when data size increases, the average runtime and I/O access of the three approaches all become larger. PLUB remains the most efficient approach while AVG becomes the slowest one when data size exceeds 100K though it has the least I/O access. This demonstrates that our derived lower bound and upper bound are very effective in improving the efficiency of query processing. The sublinear growth of runtime of our proposed approaches indicates that our approaches are applicable to large datasets.

3.5.4 Effect of Dimensionality

The dataset of the same size (100K) becomes sparser in a higher dimensional space. In other words, the object density decreases with increasing $d$. A smaller object density contributes to better performance of query processing since it becomes easier to prune unpromising objects. On the other hand, a higher $d$ also means more complicated computation and thus slows down query processing. Fig. 3.5(a) and Fig. 3.5(b) show the effect of dimensionality on runtime and I/O access of the three approaches. When $d$ is small, the impact of object density overrides that of computation, which results in decreasing runtime. When $d$ is larger than 3, the impact of computation dominates the overall query processing and leads to increasing runtime. However, even when $d = 6$, the runtime is still less than 3 seconds (1 second for PLUB). Hence, our approaches can be applied to
higher dimensions. This also indicates that PLUB is still best choice among the three approaches in terms of efficiency.

### 3.5.5 Discussion

Based on the above analyses, we conclude that PLUB achieves the best performance in efficiency because it uses both lower bound and upper bound for filtering. AVG is a better choice than PLB when $k$ is larger than 10. But when data size is larger than 100K, PLB becomes to be preferred to AVG. In general, all of our proposed approaches can answer queries in seconds while the naïve way of sequential scan may take hours or even days. The index construction time of the three approaches does not differ very much with the same data size. For example, when data size is 100K, AVG, PLB, and PLUB takes about 0.266, 0.340, and 0.345 seconds, respectively. We can also see that the time used for index construction is quite small. PLB and PLUB have the same size of R-tree and their R-trees are slightly larger than that of AVG.

### 3.6 Related Work

*Nearest neighbor search* is a fundamental problem in such diverse areas that this problem has been studied extensively and is now relatively well understood. A survey on this problem is given in [89]. As uncertainty is inherent and immerging in many applications, there are a number of studies working on nearest neighbor searching under
uncertainty \cite{22, 23, 90}, in which uncertainties are represented by probability distributions.

In \cite{22}, Ljosa et al. propose an approximate scheme to index expected distance for nearest neighbor search under the $L_1$ distance. Their result cannot be applied to our problem since we consider the exact case under the squared Euclidean distance. In \cite{23}, Agarwal et al. also consider the expected distance from a query point to an uncertain object and return expected nearest neighbor (ENN). They propose efficient algorithms for answering ENN queries under several distance functions including squared Euclidean distance. However, their discussions focus on theoretic computation of the expected Voronoi diagram as preprocessing to answer ENN queries.

On the other hand, \cite{90} considers the aspect of probability and studies probabilistic nearest neighbor (PNN) queries where the qualification probability of an object being the nearest neighbor of a query point is larger than a threshold (0 or a specified probability threshold). There is also much work studying top-$k$ probable nearest neighbor \cite{91}, superseding nearest neighbor \cite{92}, and ranking queries \cite{93}.

However, none of the above work pays particular attention to Gaussian distribution, which is a popular probability distribution in many fields of applications. In \cite{26}, Böhm et al. model the feature vector of an uncertain object using Gaussian distribution and find similar Gaussian distributions to a given query Gaussian distribution. The restriction of this work is that they assume all Gaussian distributions are probabilistically independent in each dimension, which makes it difficult to be applied to generic Gaussian distributions and limits its overall accuracy of the query result. Given a query object (a point or Gaussian distribution), the work in \cite{20} retrieves data objects represented by Gaussian distributions that are with a certain range from the query with probabilities larger than a specified threshold. They are both different from the problem we study in this work.

Finally, it is worth mentioning that nearest neighbor search has numerous variations such as reverse nearest neighbor search \cite{94}, aggregate nearest neighbor search \cite{95}, continuous nearest neighbor search \cite{96}, etc.
Chapter 3  

3.7 Summary

In this work, we considered $k$-nearest neighbor search over objects represented by Gaussian distributions based on expected distance. To support efficient query processing, we analyzed mathematically the properties of expected distance on Gaussian distribution and propose three novel approaches: AVG, PLB, and PLUB. We demonstrated the efficiency of our proposed approaches through extensive experiments and a comprehensive performance study. Among the three approaches, PLUB achieves the best efficiency while AVG is better than PLB if $k$ is large, and PLB is a better choice in the case of a large data size.
Chapter 4

Top-$k$ Similarity Search over Gaussian Distributions Based on KL-Divergence

4.1 Introduction

Probabilistic modeling, which infers probability distributions from vast amount of data for real-world applications, is being practiced in a wide range of fields from statistics, machine learning and pattern recognition \[15\] to bioinformatics and medical informatics \[97\]. The research on managing probabilistic model-based data was pioneered by Deshpande et al. \[66\], and then received considerable attention from the database research community \[8, 9, 11\]. In this work, we study the problem of processing similarity search queries over probabilistic model-based data, specifically, over objects represented by Gaussian distributions. As shown in Fig. 4.1 given a database of Gaussian distributions $G$ and a query Gaussian distribution $q$, our objective is to find top-$k$ Gaussian distributions from $G$ that are similar to $q$. In this work, we study query processing over Gaussian distributions in view of their property of being probability distributions rather than using them to represent uncertainties in Chapter 2 and Chapter 3.

Gaussian distribution, one of the most typical probability distributions, is widely used in statistics, pattern recognition, and machine learning \[15, 16\]. Research work on Gaussian
distributions has been conducted over a long period of time, including music classification [98] and search [99], and image retrieval [24, 100].

For this reason, we focus on similarity search over data modeled in Gaussian distributions, assuming that a large number of objects, represented by non-correlated Gaussian distributions are stored in the database. By non-correlated Gaussian distribution, we mean that all dimensions are independent with each other, i.e., the covariance matrix of each Gaussian distribution is diagonal. In this work, we focus on non-correlated Gaussian distributions since they are frequently used in machine learning and statistics. Hereafter, we use the term Gaussian distributions for non-correlated ones. We will report query processing methods for the general correlated case in the future work because they are very different. Given a query Gaussian distribution, our task is to retrieve from the database the Gaussian distributions that are similar to the query. The top-\(k\) Gaussian distributions with the highest similarity scores are returned as the answer to the query.

In [24], Böhmer et al. considered similarity search on feature vectors such as structural features of 2-D contours [101], time series [102] and color histograms in image databases. They represented the uncertainty of each feature vector using a non-correlated multidimensional Gaussian distribution. As discussed in [15], compared to general Gaussian distributions, the number of parameters, the storage and computational requirements can be reduced substantially by using non-correlated Gaussian distributions. In consequence, the non-correlated Gaussian distribution is often preferred in practical applications [103].

Furthermore, Gaussian mixture models (GMMs), which are linear combinations of Gaussian distributions, are known for their ability to model arbitrarily shaped distributions.
For instance, GMMs are used to model driver behaviors for driver identification in [104]. We believe that our work paves the way for similarity search over GMMs, which will be beneficial for many real world applications such as finding drivers with similar driving behaviors.

To capture the similarity between a data Gaussian distribution and a query Gaussian distribution, we choose Kullback-Leibler divergence (KL-divergence) [27], a representative measure for quantifying the similarity between two probability distributions. KL-divergence is introduced in [105], and has then become the commonest divergence measures used in practice [106].

It is well-known that KL-divergence is a non-metric measure which violates the properties of a standard distance function in metric spaces such as the Euclidean space with the Euclidean distance. Specifically, it is asymmetric and does not satisfy the triangular inequality. Hence, existing index structures based on distance functions for metric spaces like M-tree [107] cannot be employed to solve this problem.

A naïve solution is to sequentially compute the KL-divergence with the query Gaussian distribution for each Gaussian distribution in the database, and select ones with top-$k$ smallest KL-divergences. However, this method poses remarkable computing overhead and hence is not scalable to large datasets. In consequence, we employ the filter-and-refine paradigm to improve the efficiency of query processing. It first generates a set of promising candidates and filters unpromising ones without computing their similarities, and then candidate objects are refined to obtain the final results.

We propose two types of approaches utilizing the notions of rank aggregation [32] and skyline queries [33]. The first type presorts all objects in the database on their attributes and computes result objects by merging candidates from presorted lists. We modify the representative threshold algorithm (TA) [32] and propose two algorithms for efficient query processing. The second one transforms the problem to the computation of dynamic skyline queries [108]. We extend and modify the branch-and-bound skyline (BBS) algorithm [108], which is proposed to answer skyline queries, and develop a novel algorithm to solve this problem.
We note that although there have been several studies on searching in non-metric spaces [28–31], they are mainly developed for the generic class of non-metric similarity measures in discrete domains. None of them paid particular attention to the case where objects are modeled in Gaussian distributions and KL-divergence is chosen as the similarity measure. To the best of our knowledge, our work is the first study in similarity search based on KL-divergence over Gaussian distributions.

Our contributions are listed as follows.

1. We formalize the problem of top-$k$ similarity search based on KL-divergence over Gaussian distributions, and analyze mathematically the properties of KL-divergence between two Gaussian distributions.

2. We propose two types of approaches to improve the efficiency of query processing using the notion of rank aggregation and skyline queries.

3. We demonstrate the efficiency and effectiveness of our approaches through a comprehensive experimental performance study.

The rest of this chapter is organized as follows. We formally define the problem in Section 4.2. Then we analyze KL-divergence of Gaussian distributions in Section 4.3 and propose two types of approaches in Section 4.4 and Section 4.5. Experimental results and analyses are presented in Section 4.6. We review related work in Section 4.7. Finally, Section 4.8 concludes this chapter.

### 4.2 Problem Definition

#### 4.2.1 Gaussian Distribution

In the one-dimensional space, a Gaussian distribution is described by the average $\mu$ and the variance $\sigma^2$:

$$p(x) = \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left[-\frac{(x-\mu)^2}{2\sigma^2}\right]. \quad (4.1)$$
In the $d$-dimensional space, it is represented by the average vector $\mu$ and the covariance matrix $\Sigma$ [15]:

$$p(x) = \frac{1}{(2\pi)^{d/2}|\Sigma|^{1/2}} \exp\left[-\frac{1}{2}(x - \mu)^T \Sigma^{-1} (x - \mu)\right].$$

(4.2)

$|\Sigma|$ (resp. $\Sigma^{-1}$) is the determinant (resp. inverse matrix) of $\Sigma$. ($\cdot)^T$ means the transposition of ($\cdot$). In this work, we assume that a large number of objects represented by Gaussian distributions are stored in the database. For simplicity, objects represented by Gaussian distributions are called Gaussian objects, and Gaussian objects in the database are called data Gaussian objects afterwards.

### 4.2.2 Similarity Measure: KL-divergence

Given two continuous probability distributions $p_1(x)$ and $p_2(x)$, the Kullback-Leibler divergence (KL-divergence) or relative entropy [27] between them is

$$D_{KL}(p_1\|p_2) = \int_{-\infty}^{+\infty} p_1(x) \ln \frac{p_1(x)}{p_2(x)} dx.$$ 

(4.3)

In information theory, KL-divergence $D_{KL}(p_1\|p_2)$ is interpreted as a measure of the inefficiency of assuming that the distribution is $p_2$ when the true distribution is $p_1$ [27]. In other words, it measures the information lost when $p_2$ is used to approximate $p_1$. The smaller the KL-divergence is, the more similar the two probability distributions are. Accordingly, the problem of KL-divergence-based top-$k$ similarity search over Gaussian distributions (KLD-Gauss) is actually equivalent to finding top-$k$ Gaussian objects having the smallest KL-divergences with the query Gaussian object.

KL-divergence satisfies the following properties of standard distance functions: 1) non-negativity: $D_{KL}(p_1\|p_2) \geq 0$; 2) identity: $D_{KL}(p_1\|p_2) = 0$ if and only if $p_1(x) = p_2(x)$. However, it is not symmetric, i.e., $D_{KL}(p_1\|p_2) \neq D_{KL}(p_2\|p_1)$ in general. Moreover, it violates the notion of triangular inequality, namely, $D_{KL}(p_1\|p_2) + D_{KL}(p_2\|p_3) \geq D_{KL}(p_1\|p_3)$ does not necessarily hold. In other words, KL-divergence is a non-metric measure [31]. Hence, index structures designed for query processing in metric spaces such as M-tree [107] and iDistance [109] cannot be applied to accelerate similarity search based on KL-divergence.
Chapter 4  Top-k Similarity Search over Gaussian Distributions Based on KLD

As KL-divergence is asymmetric, given a data Gaussian object \( p \) and a query Gaussian object \( q \), there are two options when using it as the similarity measure between them: \( D_{\text{KL}}(p\|q) \) or \( D_{\text{KL}}(q\|p) \). It is not easy to decide which one to use, and may vary according to different applications. Both of them are common in the literature. Thus, in this work, we study both types.

The naïve approach of solving the \( KLD-Gauss \) problem is to perform a sequential scan over all objects in the database and compute their KL-divergences for each query. This approach is obviously too time-consuming and will induce intolerable computation cost for many real-world applications, especially over large scale databases. To improve the efficiency of query processing, we adopt the well-known filter-and-refine paradigm. The rationale is to avoid unnecessary computations by developing effective filtering techniques.

In this work, we propose two types of approaches for filtering. They utilize the notions of rank aggregation \[32]\ and skyline queries \[33]\, respectively. Below we first present our analysis of KL-divergence of Gaussian distributions, and then introduce our approaches.

### 4.3 KL-divergence of Gaussian Distributions

Given two one-dimensional Gaussian distributions \( p_1(x) = N(\mu_1, \sigma_1) \) and \( p_2(x) = N(\mu_2, \sigma_2) \), their KL-divergence, denoted as \( D_{\text{KL}}^1(p_1\|p_2) \), is as follows \[110\]:

\[
D_{\text{KL}}^1(p_1\|p_2) = \frac{1}{2} \left[ \frac{(\mu_1 - \mu_2)^2}{\sigma_2^2} + \frac{\sigma_1^2}{\sigma_2^2} - \ln \frac{\sigma_1^2}{\sigma_2^2} - 1 \right].
\]  

(4.4)

The two types of KL-divergence, \( D_{\text{KL}}^1(p\|q) \) and \( D_{\text{KL}}^1(q\|p) \), are referred to as \( D_{\text{KL},1} \) and \( D_{\text{KL},2} \), respectively.

In the \( d \)-dimensional space, given \( p_1(x) = N(\mu_1, \Sigma_1) \) and \( p_2(x) = N(\mu_2, \Sigma_2) \), \( D_{\text{KL}}^d(p_1\|p_2) \) is defined by

\[
D_{\text{KL}}^d(p_1\|p_2) = \frac{1}{2} \sum_{i=1}^{d} \left( \frac{(\mu_{1,i} - \mu_{2,i})^2}{\sigma_{2,i}^2} + \frac{\sigma_{1,i}^2}{\sigma_{2,i}^2} - \ln \frac{\sigma_{1,i}^2}{\sigma_{2,i}^2} - d \right),
\]  

(4.5)
where \( \mu_{1,i} \) (resp. \( \mu_{2,i} \)) is the \( i \)-th \( (1 \leq i \leq d) \) element of \( p_1 \) (resp. \( p_2 \))’s average vector \( \mu_1 \) (resp. \( \mu_2 \)). According to the independence assumption, \( \Sigma_1 \) and \( \Sigma_2 \) are diagonal matrices and their diagonal elements are denoted by \( \sigma_{1,i}^2 \) and \( \sigma_{2,i}^2 \), respectively. Obviously, \( D_{KL}^d = \sum_{i=1}^{d} D_{KL}^i \), where \( D_{KL}^i \) is the KL-divergence in the \( i \)-th dimension.

Similarly, the two types of \( D_{KL}^d \), \( D_{KL}^d(p\|q) \) and \( D_{KL}^d(q\|p) \), are denoted by \( D_{KL,1}^d \) and \( D_{KL,2}^d \), respectively. Since they are sums of the one-dimensional case, their properties of monotonicity in each dimension are the same to that of \( D_{KL,1}^1 \) and \( D_{KL,2}^1 \), which will be discussed subsequently.

### 4.3.1 \( D_{KL}^1(p\|q): D_{KL,1}^1 \)

As the smaller \( D_{KL,1}^1 \) is, the more similar \( p \) is to \( q \), we differentiate \( D_{KL,1}^1 \) on \( p \), specifically on \( \mu_p \) and \( \sigma_p^2 \), and obtain the following equations:

\[
\frac{\partial D_{KL,1}^1}{\partial \mu_p} = \frac{\mu_p - \mu_q}{\sigma_q^2} \tag{4.6}
\]

\[
\frac{\partial D_{KL,1}^1}{\partial \sigma_p^2} = \frac{\sigma_p^2 - \sigma_q^2}{\sigma_p^2 \sigma_q^2} \tag{4.7}
\]

By letting both Eq. \(4.6\) and Eq. \(4.7\) equal to 0, we derive the following property illustrated in Fig. \(4.2\). The arrows indicate decreasing directions of KL-divergence. We use \((\mu_p - \mu_q)^2\) as the horizontal axis to make the figure easy to understand.

**Property 3.** \( D_{KL,1}^1 \) is a monotonically increasing function centered at \((\mu_q, \sigma_q^2)\) and divided by \( \mu_p = \mu_q, \sigma_p^2 = \sigma_q^2 \).

1. As \( \mu_p \) increases, \( D_{KL,1}^1 \) increases monotonically when \( \mu_p > \mu_q \), and decreases monotonically when \( \mu_p < \mu_q \).

2. As \( \sigma_p^2 \) increases, \( D_{KL,1}^1 \) increases monotonically when \( \sigma_p^2 > \sigma_q^2 \), and decreases monotonically when \( \sigma_p^2 < \sigma_q^2 \).

3. \( D_{KL,1}^1 \) is minimized at \( \mu_p = \mu_q, \sigma_p^2 = \sigma_q^2 \), and its minimum is 0.
Obviously, $D_{KL1}^1$ is divisionally monotonous. The closer a point is to the center $(\mu_q, \sigma^2_q)$ and the dividing lines $\mu_p = \mu_q$ and $\sigma^2_p = \sigma^2_q$, the smaller $D_{KL1}^1$ is. In other words, smaller $|\mu_p - \mu_q|$ and $|\sigma^2_p - \sigma^2_q|$ lead to smaller $D_{KL1}^1$.

### 4.3.2 $D_{KL}^1(q||p)$: $D_{KL2}^1$

Similarly, we differentiate $D_{KL2}^1$ on $\mu_p$ and $\sigma^2_p$, and get the following equations:

\[
\frac{\partial D_{KL2}^1}{\partial \mu_p} = \frac{\mu_p - \mu_q}{\sigma^2_p} \quad (4.8)
\]
\[
\frac{\partial D_{KL2}^1}{\partial \sigma^2_p} = \frac{\sigma^2_p - \sigma^2_q - (\mu_p - \mu_q)^2}{\sigma^4_p} \quad (4.9)
\]

In the same way, by letting both Eq. (4.8) and Eq. (4.9) equal to 0, we can obtain the following property illustrated in Fig. 4.3. It differs from $D_{KL1}^1$ in that its plane is divided by $\sigma^2_p = \sigma^2_q + (\mu_p - \mu_q)^2$ instead of $\sigma^2_p = \sigma^2_q$.

**Property 4.** $D_{KL2}^1$ is a monotonically increasing function centered at $(\mu_q, \sigma^2_q)$ and divided by $\mu_p = \mu_q$, $\sigma^2_p = \sigma^2_q + (\mu_p - \mu_q)^2$.

1. As $\mu_p$ increases, $D_{KL2}^1$ increases monotonically when $\mu_p > \mu_q$, and decreases monotonically when $\mu_p < \mu_q$.

2. As $\sigma^2_p$ increases, $D_{KL2}^1$ increases monotonically when $\sigma^2_p > \sigma^2_q + (\mu_p - \mu_q)^2$, and decreases monotonically when $\sigma^2_p < \sigma^2_q + (\mu_p - \mu_q)^2$. 

(Figure 4.2: Property of $D_{KL1}^1$)  (Figure 4.3: Property of $D_{KL2}^1$)
3. $D_{KL2}^{1}$ is minimized at $\mu_p = \mu_q$, $\sigma_p^2 = \sigma_q^2 + (\mu_p - \mu_q)^2$, and its minimum is 0.

Similarly, the closer a point is to the center $(\mu_q, \sigma_q^2)$ and the dividing lines $\mu_p = \mu_q$ and $\sigma_p^2 = \sigma_q^2 + (\mu_p - \mu_q)^2$, the smaller $D_{KL2}^{1}$ is. Hence, smaller $|\mu_p - \mu_q|$ and $|\sigma_p^2 - \sigma_q^2 - (\mu_p - \mu_q)^2|$ indicate smaller $D_{KL2}^{1}$.

### 4.4 TA-based Query Processing

The problem of sequential scan lies in that, it has to compute KL-divergences of all objects in the database, even though only $k$ of them will be the answer. Based on this observation and monotonic properties of KL-divergence discussed in Section 3.3, we consider selecting a small number of promising candidate objects by presorting to avoid computing KL-divergences for unpromising objects. For example, in the one-dimensional case, we can sort the database on $\mu$ and $\sigma^2$ in advance, and only consider ones whose $\mu$ and $\sigma^2$ are close enough to that of $q$.

This is the basic idea of our first type of approaches, which utilize the notion of rank aggregation [32]. Generally speaking, we rank objects on each attribute and aggregate ones with high ranks to obtain the final top-$k$ objects. Below we use the representative threshold algorithm (TA) [32] for description. To better solve the $KLD-Gauss$ problem, we propose novel algorithms by modifying the TA algorithm. Below we first describe the TA algorithm and then introduce our proposed algorithms.

#### 4.4.1 The TA Algorithm

TA assumes that a middleware system $S$ aggregates answers to queries from various subsystems. Each subsystem $S_i$ supports two modes of data access: *sorted access* and *random access*. In the *sorted access* mode, $S$ obtains the grade of an object in the sorted list of $S_i$ by proceeding through the list sequentially from the top. On the other hand, *random access* to $S_i$ returns the corresponding grade of a given id.
For example, consider a database of images in Fig. 4.4 with two subsystems $S_{\text{color}}$ and $S_{\text{shape}}$. $S_{\text{color}}$ returns images based on their colors, and $S_{\text{shape}}$ is based on shapes. Given a query of $Q : (\text{color} = \text{"red"}) \land (\text{shape} = \text{"round"})$, $S$ merges images with high redness grades in $S_{\text{color}}$ and high roundness grades in $S_{\text{shape}}$ to obtain images satisfying the query.

Assume that $Q$ requests top-3 images based on the score function $s_Q(x) = \min\{s_{\text{red}}(x), s_{\text{round}}(x)\}$, where $s_{\text{red}}(x)$ (resp. $s_{\text{round}}(x)$) denotes the redness (resp. roundness) grade of image $x$. The left table lists top-5 images with their grades in each subsystem. Assume we retrieve one image by each sorted access. First, we obtain $\{s_{\text{red}}(x_1) = 0.96, s_{\text{round}}(x_2) = 0.95\}$. Other grades of $x_1$ and $x_2$, $s_{\text{round}}(x_1) = 0.76$ and $s_{\text{red}}(x_2) = 0.91$, can be obtained via random access. Thus, $s_Q(x_1) = 0.76$ and $s_Q(x_2) = 0.91$. The two images are both added into a result set $R = \{(x_2, 0.91), (x_1, 0.76)\}$. At the same time, the threshold $\tau$ is kept $\tau = \min\{0.96, 0.95\} = 0.95$. This is the possible best score of all images unprocessed. Once scores of images in $R$ are all no smaller than $\tau$, the algorithm stops and returns $R$.

Next, $s_{\text{red}}(x_2) = 0.91$ and $s_{\text{round}}(x_3) = 0.92$ are retrieved. Since $x_2$’s score has already been computed, we do random access only for $x_3$ and compute its score $s_Q(x_3) = 0.81$. Then $x_3$ is added into $R$ and $\tau$ is updated to 0.91. In the next step, since $x_4$ has a better score than $x_1$, we update $R = \{(x_2, 0.91), (x_4, 0.83), (x_3, 0.81)\}$ and $\tau = 0.85$. Finally, as $x_3$ and $x_4$ have already been processed, we only need to update $\tau = 0.81$. At this point, $\tau$ is no better than any of images in $R$, i.e., no image unprocessed can have a better score than that in $R$. Therefore, the algorithm terminates and returns $R$ as shown in the right table of Fig. 4.4 For ease of understanding, we associate each score with its corresponding attribute, i.e., red or round.
### 4.4.2 The Proposed Algorithms

To utilize TA, we redefine *sorted access* and *random access* for the KLD-Gauss problem. Given an object id, *random access* returns the corresponding average vector and covariance matrix. We design two types of *sorted access*. The first one retrieves $\mu_{p,j}$ or $\sigma_{p,j}^2$ ($1 \leq j \leq d$) and object id in the ascending order of $|\mu_{p,j} - \mu_{q,j}|$ or $|\sigma_{p,j}^2 - \sigma_{q,j}^2|$ (or $|\sigma_{p,j}^2 - \sigma_{q,j}^2 - (\mu_{p,j} - \mu_{q,j})^2|$, omit afterwards). The second one gives access to $D_{KL}^i$ ($1 \leq j \leq d$) and object id in the ascending order of $D_{KL}^i$. They are called CompleteTA (CTA) and PartialTA (PTA), respectively, and will be detailed subsequently.

#### 4.4.2.1 The CTA Algorithm

For CTA, we presort the database on $\mu_{p,j}$ and $\sigma_{p,j}^2$ ($1 \leq j \leq d$), and get $2d$ sorted lists. For example, in Fig. 4.5 the left table shows a list of 12 one-dimensional objects, and the right table shows their sorted orders on $\mu_i$ and $\sigma_i^2$ (called $S_\mu$ and $S_{\sigma^2}$, respectively).

In the multidimensional case, for each dimension $j$, we sort all objects on both $\mu_{i,j}$ and $\sigma_{i,j}^2$ and get $2d$ sorted lists: $(S_{\mu,1}, S_{\sigma^2,1}), \ldots, (S_{\mu,d}, S_{\sigma^2,d})$. By default, in each dimension $j$, the sorted access to each list of average returns an object $p$ with its average $\mu_{p,j}$ in the ascending order of $|\mu_{p,j} - \mu_{q,j}|$, and the sorted access to each list of variance returns another object $g$ with its variance $\sigma_{g,j}^2$ in the ascending order of $|\sigma_{g,j}^2 - \sigma_{q,j}^2|$.

<table>
<thead>
<tr>
<th>$g_i$</th>
<th>$\mu_i$</th>
<th>$\sigma_i^2$</th>
<th>Order</th>
<th>$S_\mu$</th>
<th>$S_{\sigma^2}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$g_1$</td>
<td>2</td>
<td>2.5</td>
<td>1</td>
<td>(2, $g_1$)</td>
<td>(0.2, $g_{11}$)</td>
</tr>
<tr>
<td>$g_2$</td>
<td>3.5</td>
<td>2.1</td>
<td>2</td>
<td>(3.5, $g_2$)</td>
<td>(0.5, $g_{12}$)</td>
</tr>
<tr>
<td>$g_3$</td>
<td>5</td>
<td>2.7</td>
<td>3</td>
<td>(4, $g_7$)</td>
<td>(0.7, $g_7$)</td>
</tr>
<tr>
<td>$g_4$</td>
<td>7</td>
<td>2.4</td>
<td>4</td>
<td>(5, $g_3$)</td>
<td>(0.8, $g_{10}$)</td>
</tr>
<tr>
<td>$g_5$</td>
<td>9</td>
<td>2.5</td>
<td>5</td>
<td>(6, $g_8$)</td>
<td>(1.1, $g_9$)</td>
</tr>
<tr>
<td>$g_6$</td>
<td>8.5</td>
<td>0.2</td>
<td>6</td>
<td>(6.4, $g_9$)</td>
<td>(1.2, $g_8$)</td>
</tr>
<tr>
<td>$g_7$</td>
<td>6.4</td>
<td>1.2</td>
<td>7</td>
<td>(7, $g_4$)</td>
<td>(1.8, $g_6$)</td>
</tr>
<tr>
<td>$g_8$</td>
<td>6</td>
<td>1.1</td>
<td>8</td>
<td>(8, $g_6$)</td>
<td>(2.1, $g_2$)</td>
</tr>
<tr>
<td>$g_9$</td>
<td>9.6</td>
<td>0.8</td>
<td>9</td>
<td>(8.5, $g_{11}$)</td>
<td>(2.4, $g_4$)</td>
</tr>
<tr>
<td>$g_{10}$</td>
<td>9</td>
<td>0.8</td>
<td>10</td>
<td>(9, $g_{10}$)</td>
<td>(2.5, $g_1$)</td>
</tr>
<tr>
<td>$g_{11}$</td>
<td>8.5</td>
<td>0.2</td>
<td>11</td>
<td>(9, $g_5$)</td>
<td>(2.5, $g_5$)</td>
</tr>
<tr>
<td>$g_{12}$</td>
<td>10.6</td>
<td>0.5</td>
<td>12</td>
<td>(10.6, $g_{12}$)</td>
<td>(2.7, $g_3$)</td>
</tr>
</tbody>
</table>

**Figure 4.5:** An example dataset and its sorted orders
Algorithm 5: The straightforward CTA algorithm

1: Initialize the top-$k$ list $R$ and the query $q \ (\mu_q, \sigma^2_q)$;
2: repeat
3:   for each dimension $j$ do
4:     SortedAccess$_{\text{avg}}() \rightarrow (\mu_{a,j}, g_a)$;
5:     SortedAccess$_{\text{var}}() \rightarrow \{(\sigma^2_{v,j}, g_v), (\sigma^2_{v,j}, g_v')\}$;
6:     if Any of $g_a, g_v, g_v'$ has not been accessed then
7:       Do RandomAccess and calculate its KL-divergence;
8:     Update $R$;
9:     Let $\mu_j, \overline{\sigma}^2_j, \overline{\sigma}^2_j$ be the last values accessed by SortedAccess;
10:    $\mu \leftarrow \{\mu_j | 1 \leq j \leq d\}; \overline{\sigma}^2 \leftarrow \{\sigma^2_j | 1 \leq j \leq d\}; \overline{\sigma}^2 \leftarrow \{\sigma^2_j | 1 \leq j \leq d\}$;
11:    $\tau \leftarrow \min\{\text{CalcKLD}(\mu, \overline{\sigma}^2), \text{CalcKLD}(\mu, \overline{\sigma}^2)\}$;
12: until $|R| = k$ and KL-divergences in $R$ are all no greater than $\tau$;
13: return $R$;

Algorithm 5 shows the straightforward application of the TA algorithm using the redefined random access and sorted access. The algorithm runs by dimensions. In each dimension $j$, we retrieve an object $g_a$ with its average $\mu_{a,j}$ by sorted access to the list of average. The average value of the retrieved object, $\mu_{a,j}$, is closest to that of the query, $\mu_q$, i.e., $|\mu_{a,j} - \mu_q|$ is the smallest. If there is a tie, i.e., there are two objects $g_a$ and $g_{a'}$ satisfying $|\mu_{a,j} - \mu_q| = |\mu_{a',j} - \mu_q| = \Delta$, we can break it randomly since the algorithm relies on $\tau$ and the computation of $\tau$ depends on $\Delta$ not $\mu_{a,j}$ or $\mu_{a',j}$.

On the other hand, because of the asymmetry of KL-divergence discussed in Section 4.2, each sorted access to the list of variance should return two objects in the two directions of $\sigma^2_{v,j}$: $\sigma^2_{v,j} \geq \sigma^2_{q,j}$ and $\sigma^2_{v,j} < \sigma^2_{q,j}$ (one in each direction) instead of one object to ensure correctness. We explain it using the following example. Consider a query $q$ with $\mu_q = 5, \sigma^2_q = 1$ and $k = 3$. In the first step, while we retrieve $(5, g_3)$ from $S_{\mu}$, from $S_{\sigma^2}$ both $(1.1, g_{9})$ and $(0.8, g_{10})$ need to be retrieved (in bold) to ensure correctness because we do not know which of them will have a smaller KL-divergence with respect to $q$. If we retrieve only $(1.1, g_{9})$ and find $(5, 0.8)$ has a smaller KL-divergence than that of $(5, 1.1)$ with respect to $q$ when computing $\tau$, $\tau$ will be larger than it is supposed to be and this may lead to a wrong result. In other words, we start processing from the entries in bold and continue searching in both directions.

If an object is accessed for the first time by sorted access, we obtain its average vector and covariance matrix by random access and calculate its KL-divergence (Line 6–7).
For example, we do random access for $g_3$, $g_9$ and $g_{10}$, and compute their KL-divergences ($D_{KL,1}^i$ is used for computing KL-divergences in this example).

Then we update the top-$k$ list $R$ as follows. When $|R| < k$, the object with its KL-divergence is added to $R$ directly. When $|R|$ achieves $k$, if its KL-divergence is better than any entry in $R$, we add it into $R$ and delete the entry with the largest KL-divergence so that $R$ only maintains the best $k$ objects. Meanwhile, we compute the threshold $\tau$ using the last accessed average and variance values (Line 9–11). As $\tau$ is the best KL-divergence of all objects unseen, the algorithm terminates when KL-divergences in $R$ are all no greater than $\tau$.

We show the processing steps of the example query in Fig. 4.6. Continuing with the example, we update $R = \{(g_3, 0.35), (g_9, 0.98), (g_{10}, 8.01)\}$. At the same time, we compute KL-divergences of $(5, 1.1)$ and $(5, 0.8)$ with respect to the query $(5, 1)$, and update $\tau$ as the smaller one, which is 0.002.

In the second step, $(4, g_7)$ from $S_{\mu}$, $(1.2, g_8)$ and $(0.7, g_7)$ from $S_{\sigma^2}$, are retrieved. Since KL-divergences of $g_7$ and $g_8$ are smaller than that of $g_9$ and $g_{10}$, we update $R$ as $\{(g_3, 0.35), (g_8, 0.508), (g_7, 0.53)\}$ and $\tau = 0.508$. In the last step, we retrieve $(6, g_8)$ from $S_{\mu}$, $(1.8, g_6)$ and $(0.5, g_{12})$ from $S_{\sigma^2}$. $R$ stays the same, but $\tau$ is updated to 0.597. Finally, as all the objects in $R$ have no greater KL-divergences than $\tau$, we stop searching and return $R$.

In each step, as the straightforward algorithm retrieves two objects with respect to the variance in a brute-force way, it tends to do many unnecessary accesses. We avoid them...
by considering the priority of the object in each direction and access only one in each step. We derive the following lemma to guide the algorithm (see the proof in C.1).

**Lemma 4.1.**

1. Assume $|\mu_{p,j} - \mu_{q,j}| = |\mu_{p',j} - \mu_{q,j}|$ ($1 \leq j \leq d$) and $\sigma^2_{p,m} = \sigma^2_{p',m}$ ($1 \leq m \leq d, m \neq j$). If $\sigma^2_{p,j} > \sigma^2_{q,j}$, $\sigma^2_{p',j} < \sigma^2_{q,j}$, and $(\sigma^2_{p,j} - \sigma^2_{q,j}) \leq (\sigma^2_{q,j} - \sigma^2_{p,j})$, then $D_{KL}(p||q) < D_{KL}(p'||q)$.

2. Assume $|\mu_{p,j} - \mu_{q,j}| = |\mu_{p',j} - \mu_{q,j}|$ ($1 \leq j \leq d$) and $\sigma^2_{p,m} = \sigma^2_{p',m}$ ($1 \leq m \leq d, m \neq j$). If $\sigma^2_{p,j} > \sigma^2_{q,j} + (\mu_{p,j} - \mu_{q,j})^2$, $\sigma^2_{p',j} < \sigma^2_{q,j} + (\mu_{p',j} - \mu_{q,j})^2$, and $(\sigma^2_{p,j} - \sigma^2_{q,j} - (\mu_{p,j} - \mu_{q,j})^2) \leq (\sigma^2_{q,j} + (\mu_{p',j} - \mu_{q,j})^2 - \sigma^2_{p',j})$, then $D_{KL}(q||p) < D_{KL}(q||p')$.

Based on Lemma 4.1, during the bidirectional search over the variance, when an object $p$ with $\sigma^2_{p,j} > \sigma^2_{q,j}$ is obtained, we only need to consider another object $p'$ with $\sigma^2_{p',j} < \sigma^2_{q,j}$, if $\sigma^2_{p',j}$ is nearer to $\sigma^2_{q,j}$ (or $\sigma^2_{q,j} + (\mu_{p',j} - \mu_{q,j})^2$) than that of $p$. For example, in the first step, in $S_{\sigma^2}$ when comparing $(1.1, g_9)$ with $(0.8, g_{10})$, since $(1.1 - 1) \leq (1 - 0.8)$, we only retrieve $g_9$ and delay the retrieval of $g_{10}$. In other cases, we compare their KL-divergences using the current average value $\mu_j$ obtained, i.e., KL-divergences of $(\mu_j, \sigma^2_{p,j})$ and $(\mu_j, \sigma^2_{p',j})$, and select the one with a smaller KL-divergence with respect to $q$.

We show the running steps of the improved CTA algorithm in Fig. 4.7. At first, we retrieve $(5, g_3)$ from $S_{\mu}$. Meanwhile, we retrieve $(1.1, g_9)$ from $S_{\sigma^2}$ based on Lemma 4.1 as explained above. Accordingly, we update $R$ and $\tau$. Then we continue to retrieve $(4, g_7)$ from $S_{\mu}$. Since $(1.2 - 1) \leq (1 - 0.8)$, we retrieve $(1.2, g_8)$ from $S_{\sigma^2}$ based on Lemma 4.1 and update $R$ and $\tau$ accordingly. In the third step, when comparing $(1.8, g_6)$ with $(0.8, g_{10})$ in $S_{\sigma^2}$, because they do not satisfy Lemma 4.1, we compute their KL-divergences using the current average value $\mu_8 = 6$. In other words, we compare KL-divergences of $(6, 1.8)$ and $(6, 0.8)$ with respect to $(5, 1)$. Then $g_{10}$ is selected. As we can see, while the straightforward CTA algorithm accessed 7 objects, the improved one only retrieved 5 objects with one additional step.
4.4.2.2 The PTA Algorithm

For PTA, in each dimension we construct a two-dimensional R-tree of all data Gaussian objects. When a query \( q \) comes, by using the skyline-based approach described in Section 4.5, we can obtain the top objects with the smallest \( D_{KL}^i \) with respect to \( q \) in each dimension \( i \) (detailed in Section 4.5.6). The PTA algorithm is similar to the CTA algorithm, except that we retrieve objects based on \( D_{KL}^i \) instead of \( \mu_i \) and \( \sigma_i^2 \) with respect to \( q \), and \( \tau \) is calculated by \( \tau = \sum_{i=1}^{d} D_{KL}^i \).

Next, we introduce another approach, which transforms the \( KLD-Gauss \) problem to dynamic skyline queries \[108\] by using the notion of skyline queries \[33\].

4.5 Skyline-based Query Processing

In this section, by utilizing the properties of KL-divergence and Gaussian distributions, we propose another approach for solving the \( KLD-Gauss \) problem. We extend and modify the BBS (Branch-and-Bound Skyline) algorithm \[108\], which is proposed to answer skyline queries. Below we first introduce the concept of skyline queries and the BBS algorithm, and then describe our extensions and the proposed algorithm.

4.5.1 Skyline Queries and Dynamic Skyline Queries

Given a dataset \( D \), a skyline query \[33\] returns all the objects not dominated by others within \( D \). \( p_i \) is said to be dominated by \( p_j \), if \( p_j \) is better than \( p_i \) on at least one attribute, and better than or equals to \( p_i \) on other attributes. A common example is, given a list of hotels with prices and distances from a beach, to find ones having the lowest prices and the shortest distances. A hotel with $200 and 1 km from the beach is preferable to (dominate) the one of $250 and 1.5 km.

A dynamic skyline query, a variation of a skyline query, returns all the objects that are not dynamically dominated with respect to a set of dimension functions \( f_1, f_2, ..., f_m \) specified by the query \[108\]. Each function \( f_i \) takes as parameters a subset of the \( n \) attributes
in the original $n$-dimensional space. The objective is to return the skyline in the new $m$-dimensional space defined by $f_1, f_2, ..., f_m$.

Continuing with the hotel example, assume for each hotel we store its $x$ and $y$ coordinates and price $c$ (3-dimensional) information in the database. Dynamic skyline queries can be used to retrieve hotels that are closest to a specified location $(x_u, y_u)$ and price $c_u$. In other words, closeness functions are defined by $f_1 = \sqrt{(x - x_u)^2 + (y - y_u)^2}$, and $f_2 = |c - c_u|$. Note that $(x_u, y_u)$ and $c_u$ normally vary with queries.

In [108], Papadias et al. proposed the BBS algorithm for processing skyline queries. They proved that BBS is I/O optimal; that is, it accesses the least number of disk pages among all algorithms based on R-trees. Hence, the following discussion concentrates on this technique.

### 4.5.2 The BBS Algorithm

Consider the task of computing the skyline of the example dataset in Fig. 4.5, i.e., finding objects with the smallest averages and variances. According to BBS, we construct an R-tree to index all objects. Each Gaussian object is inserted into the R-tree as a two-dimensional point $(\mu_i, \sigma^2_i)$. Its image and hierarchical structure are shown in Fig. 4.8 and Fig. 4.9. Each group of objects, i.e., an R-tree node, is represented by an MBR (Minimum Bounding Rectangle).

A priority queue $Q$ is employed to maintain entries (R-tree nodes or Gaussian objects) to be accessed in the ascending order of $\text{mindist}$. The $\text{mindist}$ of an entry, is the smallest cityblock ($L_1$) distance of its MBR to a reference point. For example, the $\text{mindist}$ of $N_1$ to the origin $O$ is calculated by summing up the length of $OA$ and $AB$, where $B$ is the lower-left point of $N_1$, and $A$ is the projection of $B$ on the $\mu_p$-axis. The $\text{mindist}$ of a Gaussian object to $O$, e.g., $g_1 = (2, 2.5)$, is simply its $L_1$ distance to $O$, i.e., $2 + 2.5 = 4.5$.

We use the example in Fig. 4.8 to illustrate the algorithm. After expanding the root node, entries in $Q$ are $\{(N_1, 3.8), (N_2, 4.2)\}$. Then we expand $N_1$ and insert $N_{11}, N_{12}$ into $Q$. $Q$ becomes $\{(N_{11}, 4.1), (N_2, 4.2), (N_{12}, 8.8)\}$. When expanding $N_{11}$, since $g_3$ is dominated...
by $g_1$ (and $g_2$), it is rejected. After expanding $N_2$, $Q = \{(g_1, 4.5), (N_{21}, 4.7), (g_2, 5.6), (N_{22}, 8.7), (N_{12}, 8.8)\}$.

Next $g_1$ is added into a candidate set $S$ as the first skyline object. Since $N_{21}$ is not dominated by $g_1$, it is expanded and $g_7$ is inserted into $Q$. Then $Q = \{(g_7, 4.7), (g_2, 5.6), (N_{22}, 8.7), (N_{12}, 8.8)\}$. $g_7$ and then $g_2$ are both added into $S$ because $g_7$ is not dominated by $g_1$, and $g_2$ is not dominated by $g_7$ and $g_1$. Subsequently, expanding $N_{22}$ leads to another candidate $g_{11}$. The last entry $N_{12}$ will not be expanded as it is dominated by $g_7$. Finally, $S = \{g_1, g_7, g_2, g_{11}\}$ is returned as the skyline result.

BBS can be applied to compute dynamic skylines by expanding entries in $Q$ according to $\text{mindist}$ in the dynamic space \cite{108}. In other words, we compute $\text{mindist}$ with respect to the query object $q$ instead of the origin $O$. Dynamic skylines can be computed in the same way except that the $\text{mindist}$ of each entry in $Q$ will be changed (each $\text{mindist}$ is computed on-the-fly when the entry is considered for the first time). Assuming $f_1 = |\mu_p - \mu_q|$ and $f_2 = |\sigma^2_p - \sigma^2_q|$, the dynamic skyline result of the query $q = (5, 1)$ in Fig. 4.8 is $\{g_3, g_8, g_9\}$.

### 4.5.3 Transformation and Extension

According to Section 4.3, the closer $\mu_p$ is to $\mu_q$, $\sigma^2_p$ is to $\sigma^2_q$, the smaller KL-divergence is. This is analogous to a dynamic skyline query by assuming $f_1 = |\mu_p - \mu_q|$, $f_2 = |\sigma^2_p - \sigma^2_q|$ (or
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\[ f_2 = |\sigma_p^2 - \sigma_q^2 - (\mu_p - \mu_q)^2| \]

Hence, we transform the KLD-Gauss problem to computing the dynamic skyline with respect to \( q \).

In [108], BBS is applied directly to compute dynamic skylines. However, we note that since KL-divergence is asymmetric over \( \sigma_i^2 \), the two subspaces divided by \( \sigma_p^2 = \sigma_q^2 \) (or \( \sigma_p^2 = \sigma_q^2 + (\mu_p - \mu_q)^2 \)) should be treated separately when we use BBS. In each dimension, we need to maintain two priority queues, and merge two result sets to obtain the final top-\( k \) objects. This is obviously inefficient and would incur extra computation cost.

Note that a straightforward method is to use iterative skyline computation according to the property that the top-1 object, based on any monotone ranking function, must be one of the skyline objects [111]. After deleting the top-1 object from our consideration, we recompute dynamic skyline objects. Then we select the top-2 object among them. Top-3, 4, \ldots, \( k \) objects are computed in a similar way. This method would also lead to heavy cost.

The two concerns motivate us to propose a more efficient approach. We develop our ideas by extending the BBS algorithm. For ease of presentation, we describe our approach using \( D_{KL}^1 \) and present necessary modifications in Section 4.5.5.

Because of the asymmetry of KL-divergence, we can only determine the dominance relationship between an object and each entry in the same subspace where the object locates, and cannot conclude whether it dominates any entry in the other subspace directly. For example, in Fig. 4.8 we can directly determine that \( g_8 \) dominates \( N_{12} \). However, we do not know whether \( g_8 \) dominates \( g_7 \) or \( N_{22} \). As a result, the dynamic skyline in each subspace has to be computed separately.

In order to reduce the cost caused by separate subspace computation, we consider enhancing the filtering power of each object from the subspace where it locates to the other subspace. If the dominance relationship between the two subspaces can be determined, we only need to maintain one priority queue and compute the two dynamic skylines together. To this end, for each point in one subspace, we find its counterpoint in the other subspace so that it can be used for filtering in that subspace.
This idea is confirmed by the properties of KL-divergence discussed in Section 4.3.3. Since KL-divergence shows monotonicity in both sides of $\sigma_p^2 = \sigma_q^2$, for each point $g$ in one subspace there must be one and only one corresponding point $g'$ that satisfies $\mu_{g'} = \mu_g$ and $D_{KL}(g'\|q) = D_{KL}(g\|q)$. We call $g'$ the equal-KLD point of $g$. The two equations produce $\sigma_{g'}^2 - \sigma_q^2 \ln \sigma_{g'}^2 = \sigma_g^2 - \sigma_q^2 \ln \sigma_g^2$. Because of its complexity, there is no analytical solution for $\sigma_{g'}^2$. We compute $\sigma_{g'}^2$ numerically by the Newton method and use its approximation (a slightly larger value to ensure correctness) instead. For example, the equal-KLD point of $g_8 = (6, 1.2)$ is $g'_8 = (6, 0.82)$, which can be used to filter entries such as $N_{22}$ additionally (i.e., $g'_8$ dominates $N_{22}$).

We formally define the equal-KLD point as follows.

**Definition 4.2 (equal-KLD point).** Given a Gaussian point $g$, $g'$ is the equal-KLD point of $g$, if $g'$ satisfies $D_{KL}(g'\|q) = D_{KL}(g\|q)$.

Accordingly, we extend the definition of dynamically dominate to dynamically KLD-dominate as follows.

**Definition 4.3 (dynamically KLD-dominate).** Given a Gaussian point $g$ and an entry $e$, $g$ dynamically KLD-dominates $e$, if $g$'s equal-KLD point $g'$ dynamically dominates $e$.

For example, by computing $g_8$'s equal-KLD point $g'_8 = (6.01, 1.1)$, we can conclude that $g_8$ dynamically KLD-dominates $g_9 = (6.4, 1.1)$ since $g'_8$ dynamically dominates $g_9$. As $g$ is an equal-KLD point of itself, we can equally say $g$ dynamically KLD-dominates $e$, if $g$ dynamically dominates $e$. The relationship of $g$ dynamically KLD-dominates $e$ guarantees that the KL-divergence of $g$ is smaller than that of any object within $e$.

To reduce the overhead of iterative skyline computation, we modify the way of dominance checking that the BBS algorithm does. We sort all candidates in the ascending order of KL-divergence. For each entry in the priority queue, we check the dominance relationship from the $k$-th candidate to the last one instead of checking all candidates as BBS does. If all entries are dynamically KLD-dominated by these “inferior” (from $k$ to the last) candidates, the “superior” top-($k-1$) and the first “inferior” candidates will be the final top-$k$ result.
Algorithm 6 Skyline-based query processing algorithm

1: procedure KLD_QUERY_SKY(R-tree, q, k)  
2: \hspace{1em} Q \leftarrow (0, \text{Root}); \quad \triangleright \text{Initialize } Q \text{ with the Root of R-tree}  
3: \hspace{1em} S \leftarrow \emptyset; \quad \triangleright \text{Initialize the candidate set}  
4: \hspace{1em} \text{while } Q \text{ is not empty do}  
5: \hspace{2em} e \leftarrow Q.\text{top}(); Q.\text{pop}();  
6: \hspace{2em} \text{Check } e \text{ against } S;  
7: \hspace{2em} \text{if } e \text{ is not dynamically KLD-dominated by } S \text{ then}  
8: \hspace{3em} \text{if } e \text{ is a Gaussian object } g \text{ then}  
9: \hspace{4em} \text{Add } g \text{ and its KL-divergence into } S;  
10: \hspace{3em} \text{else} \quad \triangleright e \text{ is an R-tree node } N  
11: \hspace{4em} \text{for each child } N_i \text{ do}  
12: \hspace{5em} \text{Check } N_i \text{ against } S;  
13: \hspace{5em} \text{if } N_i \text{ is not dynamically KLD-dominated by } S \text{ then}  
14: \hspace{6em} Q.\text{push}(N_i, \text{mindist}(N_i));  
15: \hspace{1em} \text{return } \text{top-}k \text{ of } S;  

4.5.4 The Proposed Algorithm: SKY

Based on the discussion above, we propose the SKY algorithm shown in Algorithm 6. An R-tree is employed to index averages and variances of all Gaussian objects in the database. Given the constructed R-tree, a query Gaussian object q, and a constant k, the algorithm returns the top-k Gaussian objects having the smallest KL-divergences with q.

The algorithm begins from the root node and continues until all entries in the priority queue Q are processed. When checking the top entry e of Q against the candidate set S (Line 6), if \(|S| < k\), we add it to S if it is an Gaussian object, and expand it in the case of an R-tree node. Otherwise (i.e., \(|S| \geq k\)), we sort S in the ascending order of KL-divergence, and compare e against “inferior” candidates in S, i.e., from the k-th candidate until the last one. In this way, we can avoid the expensive iterative dynamic skyline computation. We reject e if it is dynamically KLD-dominated by S. If not, it is either added into S with its divergence (when e is a Gaussian object) or expanded (when e is an R-tree node). In the expansion of an R-tree node N, we check each of its child nodes N_i against S and insert N_i to Q if N_i is not dynamically KLD-dominated by S. Finally, the top-k candidates of S will become the result.

Another problem is that, there still will be quite a long list of entries in Q waiting for dominance checking. In order to prune non-promising entries, for each “inferior” candidate
According to Lemma 4.4, we only need to consider entries with \( \text{mindist} < \text{mmdist} \) so that the searching process can be finished early. We note that this filtering technique only works for \( D_{\text{KL}}^1 \). In other cases, all entries in \( Q \) have to be processed.

We use the example in Fig. 4.8 to illustrate our algorithm in the case of \( D_{\text{KL}}^1 \). Assume \( k = 3 \). After expanding the root node, \( Q = \{(N_2, 0), (N_1, 0.8)\} \). Then we expand \( N_2 \) and \( Q = \{(N_2, 0), (N_1, 0.8), (N_{22}, 3.7)\} \). Next, \( N_{21} \) is expanded and \( g_7, g_8, g_9 \) are inserted into \( Q \). Entries in \( Q \) are \( \{(N_1, 0.8), (g_8, 1.2), (g_7, 1.3), (g_9, 1.5), (N_{22}, 3.7)\} \).

After expanding \( N_1 \), \( Q \) is \( \{(N_{11}, 1.1), (g_8, 1.2), (g_7, 1.3), (g_9, 1.5), (N_{12}, 2.8), (N_{22}, 3.7)\} \). Then \( N_{11} \)'s child objects \( g_1, g_2, g_3 \) are inserted into \( Q \). Thus, \( Q = \{(g_8, 1.2), (g_7, 1.3), (g_9, 1.5), (g_3, 1.7), (g_2, 2.6), (N_{12}, 2.8), (N_{22}, 3.7), (g_1, 4.5)\} \).

The top three entries \( g_8, g_7, g_9 \) with KL-divergences are added into \( S \) successively and \( S = \{(g_8, 0.51), (g_7, 0.53), (g_9, 0.98)\} \). At the same time, we compute the mmdist of \( g_9 \), which is 4.03, and keep \( \delta = 4.03 \). Next, since \( g_3 \) is not dynamically KLD-dominated by \( g_9 \), it is inserted into \( S \) and \( S = \{(g_3, 0.35), (g_8, 0.51), (g_7, 0.53), (g_9, 0.98)\} \). The mmdist of \( g_7 \) is 2.40 < 4.03. Thus, \( \delta \) is updated to 2.40. Since the mindist of \( g_2 \) is larger than \( \delta \), the algorithm stops and returns \( \{(g_3, 0.35), (g_8, 0.51), (g_7, 0.53)\} \).
4.5.5 Extending the Skyline-based Approach

The case of $D_{KL}^1$ can be processed similarly except the filtering technique based on $mmdist$ cannot be applied. In the multi-dimensional case, the algorithm is the same to the one shown in Algorithm 6. A 2d-dimensional R-tree is constructed to index d-dimensional Gaussian objects in the database. The following definitions and computations will replace their counterparts in the one-dimensional case: (1) $mindist = \sum_{i=1}^{d}(|\mu_{p,i} - \mu_{q,i}| + |\sigma^2_{p,i} - \sigma^2_{q,i}|)$ (2) Compute equal-KLD point based on $D_{KL1}^d$ or $D_{KL2}^d$. (3) An object $g$ dynamically KLD-dominates an entry $e$, if $g$’s equal-KLD point $g'$ dynamically dominates $e$ in all dimensions.

4.5.6 Application to the PTA Algorithm

As discussed in Section 4.4.2.2, in each dimension PTA accesses the top objects with the smallest $D_{KL}^d$ using the skyline-based method SKY. Since PTA performs multiple sorted accesses in each dimension, we need to maintain all dominated entries instead of rejecting them as SKY does. We associate each “inferior” candidate with all entries dominated by it, and release them for further processing when this candidate becomes “superior” in the next sorted access.

We use the following example two-dimensional dataset shown in Table 4.1 to illustrate the PTA algorithm. We construct an R-tree in each dimension. Their R-tree images in the first dimension and the second dimension are shown in Fig. 4.10 and Fig. 4.11, respectively.

Consider the same query $q = (5.0, 2.0; 6.0, 1.5)$ with $k = 2$. Assume that we retrieve two objects in each dimension. At first, in the first dimension after expanding the root

<table>
<thead>
<tr>
<th>$g_i$</th>
<th>$\mu_{i,1}$</th>
<th>$\sigma^2_{i,1}$</th>
<th>$\mu_{i,2}$</th>
<th>$\sigma^2_{i,2}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$g_1$</td>
<td>6.0</td>
<td>0.8</td>
<td>5.8</td>
<td>2.0</td>
</tr>
<tr>
<td>$g_2$</td>
<td>6.0</td>
<td>3.0</td>
<td>6.4</td>
<td>1.6</td>
</tr>
<tr>
<td>$g_3$</td>
<td>8.2</td>
<td>1.2</td>
<td>3.1</td>
<td>1.6</td>
</tr>
<tr>
<td>$g_4$</td>
<td>2.5</td>
<td>3.4</td>
<td>4.7</td>
<td>0.9</td>
</tr>
</tbody>
</table>

Table 4.1: A two-dimensional example dataset
same way and obtain node in Fig. 4.10, we obtain $Q_1 = \{(N_1, 1.0), (N_2, 1.8)\}$ where $Q_1$ is a priority queue maintaining entries (R-tree nodes and Gaussian objects) to be processed in the ascending order of their mindists from $q$. Then we expand $N_1$ and insert its two child Gaussian objects, $g_2$ and $g_4$, into $Q_1 = \{(N_2, 1.8), (g_2, 2.0), (g_4, 3.9)\}$. Next, we expand $N_2$ in the same way and obtain $Q_1 = \{(g_2, 2.0), (g_1, 2.2), (g_4, 3.9), (g_3, 4.0)\}$. The next entries, $g_2$ and $g_1$, are added into the candidate set $S_1$ successively with their KL-divergences in this dimension, $S_1 = \{(g_2, 0.297), (g_1, 0.408)\}$. At the same time, we compute the mmdist of $g_1$ and assign it to $\delta_1 = 3.904$. Since the mindist of the next entry $g_4$ is smaller than $\delta_1$, i.e., $3.9 < 3.904$, we continue to check $g_4$. When checking it against $S_1$, we find that $g_4$ is dynamically KLD-dominated by $g_1$. Hence, we add it to the dominating list of $g_1$, and obtain $S_1 = \{(g_2, 0.297), (g_1, 0.408), ((g_4, 3.9))\}$. At this time, we have $Q_1 = \{(g_3, 4.0)\}$. Since the mindist of the next entry $g_3$ is larger than $\delta_1$, based on Lemma 4.4 we stop searching and return $g_2$ and $g_1$ with the second smallest $D_{KL}^1 = 0.408$.

In the second dimension, we calculate the smallest $D_{KL}^2$ in the same way as $D_{KL}^1$ in the first dimension. After expanding the root node in Fig. 4.11 we obtain $Q_2 = \{(N_2, 0.1), (N_1, 1.3)\}$. Then we expand $N_2$ and obtain $Q_2 = \{(g_2, 0.5), (g_1, 0.7), (N_1, 1.3)\}$. Next, we add $g_2$ and $g_1$ into $S_2$ successively with their KL-divergences in this dimension, $S_2 = \{(g_1, 0.036), (g_2, 0.054)\}$. At the same time, we compute the mmdist of $g_2$ and assign it to $\delta_2 = 0.897$. Since the mindist of the next entry in $Q_2 = \{(N_1, 1.3)\}$ is larger than $\delta_2$, again based on Lemma 4.4 we stop searching and return $g_1$ and $g_2$ with the second smallest $D_{KL}^2 = 0.054$.

What follows is that the algorithm will compute the overall KL-divergences of $g_2$ and $g_1$ in the two dimensions (0.35 and 0.44), respectively, and update $\tau = D_{KL}^1 + D_{KL}^2 = 0.462$. Since the second smallest KL-divergence 0.44 is larger than $\tau$, we first release $(g_4, 3.9)$
from $S_1$ to $Q_1$ and delete the entries of $g_1$ and $g_2$ from $S_1$ and $S_2$, and then repeat the same process in the next round until the $k$th smallest KL-divergence is no larger than $\tau$.

### 4.6 Experiments

#### 4.6.1 Experimental Setup

We generate both data Gaussian distributions and query Gaussian distributions randomly under the same setting. Each average value is generated from $(0, 1000)$, and each variance value is generated from $(0, 100)$. The parameters tested in our experiments and their default values (in bold) are summarized in Table 4.2. To test the effect of data distribution, we also generate three datasets with independent, correlated, and anti-correlated distributions, respectively, using the standard skyline data generator in [33].

We compare our TA-based algorithms CTA, PTA and skyline-based algorithm SKY, with the sequential scan methods Scan and the extended BBS algorithm BBS. In each dimension $i$ ($1 \leq i \leq d$), BBS processes objects in the two subspaces divided by $\sigma_{p,i}^2 = \sigma_{q,i}^2$ (or $\sigma_{p,i}^2 = \sigma_{q,i}^2 + (\mu_{p,i} - \mu_{q,i})^2$) separately. In other word, it maintains $2^d$ priority queues and compute each top object by merging candidate objects from these priority queues.

During the preprocessing, we build a $2d$-dimensional R-tree for SKY and BBS. Each Gaussian distribution is inserted as a $2d$-dimensional point, consisting of the $d$-dimensional average vector and $d$-dimensional variance vector. For PTA, we construct $d$ two-dimensional R-trees. Each two-dimensional R-tree maintains the average and variance value in the $i$-th dimension ($1 \leq i \leq d$), and provides sorted access to the ranked $D_{kL}$ and object id.
(see Section 4.4). Random access is supported by an Gaussian object list in the order of object id. All R-trees and lists are stored in the secondary memory.

In each experiment, we run 100 queries and use the average runtime for performance evaluation. We do not consider I/O access because the system tends to load indices of small size into main memory upon reading and the runtime is mainly spent on CPU. All experiments are conducted using a workstation with Intel Xeon(R) CPU E3-1241 v3 (3.50GHz), RAM 16GB, running Ubuntu 12.10 LTS.

### 4.6.2 Performance Evaluation

We first compare SKY with BBS. While SKY takes about 0.02 seconds using both the first type and the second type of KL-divergence (called KLD1 and KLD2 afterwards), BBS takes 2.45 seconds for KLD1, and 0.23 seconds for KLD2. This demonstrates the effectiveness of our extension and modification to the basic BBS algorithm. Because of the uneven property of KLD2 shown in Fig. [4.3], most of objects are assigned to the same priority queue. Consequently, BBS spends less time using KLD2 than that using KLD1. We leave BBS out of our consideration in the following experiments because of its clear inefficiency.

#### 4.6.2.1 Effect of $k$

Intuitively, a larger $k$ incurs more cost, since retrieving more objects potentially leads to more computations. The constantly increasing runtime of PTA, CTA and SKY shown in Fig. [4.12] follows this intuition. Scan is an exception of this intuition since it computes KL-divergence for all data Gaussian objects regardless of $k$. CTA and SKY outperform Scan over all $k$, and PTA performs better than Scan when $k < 100$ for KLD1 ($k < 60$ for KLD2). The difference is that PTA displays the most significant increase while the runtime of CTA rises more slowly, and SKY still keeps the low runtime as $k$ increases. Even when $k$ is 100, the runtime of SKY is only 12% of that of Scan.
That’s because the TA-based approaches, CTA and PTA, retrieve many objects iteratively based on their partial information in each dimension (average, variance or KL-divergence), and they need to retrieve much more objects with a larger $k$. On the other hand, SKY retrieves less objects based on their full information in all dimensions and thus costs less runtime. Especially, PTA needs to maintain all intermediate entries and compute the dynamic skyline to retrieve candidate objects in each dimension. Thus, it takes even more runtime than CTA, which retrieves candidates more easily by sorted access to presorted lists.

4.6.2.2 Effect of Data Size

Figure 4.13 shows the scalability of the four approaches. All of them consume more runtime with a larger data size. Scan exhibits the worst scalability to large scale datasets, while SKY demonstrates the best. The performance of PTA and CTA is between them with that CTA performs better.

When the data size increases, the density of objects in the space becomes larger. This means that there will be more objects with similar averages, variances or KL-divergences in one dimension but different in other dimensions. As a result, PTA and CTA suffer more from ineffective retrievals and thus perform worse than SKY, which has better scalability due to its more effective retrieval by R-tree. Especially, SKY and CTA outperform Scan when the data size is larger than 10K and are much better with the increasing data size.
4.6.2.3 Effect of Dimensionality

As shown in Fig. 4.14, dimensionality affects the proposed approaches greatly, while it has little effect on Scan. Since PTA and SKY both utilize R-trees for indexing and the performance of R-tree degrades with the increasing dimension $d$, their runtime rises very fast. The runtime of PTA rises even faster since in each dimension it retrieves objects based on partial information and this results in much more ineffective retrievals, i.e., it retrieves much more unpromising objects. CTA spends more runtime with increasing $d$ due to the similar reason as PTA that it does much more ineffective retrievals.

When $d$ is larger than 3, they deteriorate significantly and are defeated by Scan. This indicates that our proposed approaches are more efficient in low dimensions less than 4. Moreover, their performance will be improved with a larger data size. For example, as shown in Fig. 4.15 when $d = 3$, the advantage of CTA and SKY over Scan is more obvious when the data size increases from 1000K to 10000K.
4.6.2.4 Effect of Data Distribution

In Fig. 4.16, we show the effect of data distribution on the four approaches. Generally, the runtime of all approaches does not vary greatly over the three different distributions. CTA shows the most apparent decrease over correlated and anti-correlated distributions since it depends directly on the average or variance of each object in each dimension. Correlations mean that objects are concentrated in the center and thus decrease the number of ineffective retrieval on average. It is clear that SKY has the best performance. This demonstrates the capability of our proposed approaches over different data distributions.

4.6.2.5 Index Construction

As preprocessing, we build R-trees and lists to support efficient query processing. In each experiment, we build a $2d$-dimensional R-tree for SKY and BBS, and $d$ two-dimensional R-trees for PTA. When using the default dataset ($d = 2$, data size = 1000K), the index construction time is about 5 seconds for building a four-dimensional R-tree, and is about
7 seconds for building 2 two-dimensional R-trees. The lists include an object list for random access in CTA and PTA, and $d$ sorted lists for sorted access in CTA and PTA. The time for building these two kinds of lists using the default dataset is about 1 second and 9 seconds, respectively. When the data size or $d$ varies, the construction time varies proportionally. In view of the performance improvement shown in previous sections, the preprocessing cost of our proposed approaches is rather low.

4.7 Related Work

In this work, we proposed query processing approaches for top-$k$ similarity search over Gaussian distributions based on KL-divergence, a non-metric similarity measure. Skopal et al. [31] surveyed the domains employing non-metric functions for similarity search, and methods for efficient and effective non-metric similarity search. To improve the searching efficiency, a class of approaches adopt the indexing strategy based on analytic properties. They analyze the properties of a specific similarity function, and develop special access methods for that function. Another advantage is their ability to support both exact and approximate search. Our work falls into this category and supports exact search.

Other approaches are based on statistical methods. For the efficiency, they perform costly preprocessing by suitably clustering or transforming the original database into another space such as the metric space based on the distance distribution statistics, so that existing metric indexing methods can be employed [28, 29, 31]. One drawback is that they cannot provide exact results. Furthermore, expensive preprocessing is often needed prior to the indexing itself. On the contrary, our proposed approaches have low preprocessing cost and support exact similarity search.

As a general class of similarity measures including KL-divergence, Bregman divergence has also led to a stream of research work to develop various algorithms. For example, [112] proposed clustering approaches with Bregman divergence. Zhang et al. [30] developed a prune-and-refine-based searching method for Bregman divergence, which is close to our work, but it works only for discrete probability distributions (described by
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In finite domains. Moreover, they considered only the first type of asymmetric divergences.

In [24], an index structure called Gauss-tree was proposed for similarity search over multivariate Gaussian distributions. They also assumed non-correlated Gaussian distributions. Although their problem is very similar to ours, they defined a different similarity measure as follows. For a $d$-dimensional Gaussian distribution $g$, they used $N_{\mu_{g,j},\sigma_{g,j}}(x_j)$ to represent its probability density function in each dimension $j$, which is a one-dimensional Gaussian distribution with two parameters, the average $\mu_{g,j}$ and the variance $\sigma_{g,j}^2$. Given a database $DB$ and a query Gaussian distribution $q$, the similarity of $g$ in $DB$ and $q$ is defined as:

$$P(g|q) = \frac{p(q|g)}{\sum_{w \in DB} p(q|w)}$$

(4.10)

where

$$p(q|g) = \prod_{j=1}^{d} \int_{-\infty}^{+\infty} N_{\mu_{q,j},\sigma_{q,j}}(x_j) \cdot N_{\mu_{g,j},\sigma_{g,j}}(x_j) dx$$

$$= \prod_{j=1}^{d} N_{\mu_{q,j},\sigma_{q,j}+\sigma_{g,j}}(\mu_{g,j})$$

(4.11)

Here, $p(q|g)$ represents the probability density for observing $q$ under the condition that we already observed $g$. The conditions that maximize Eq. (4.11) are $\mu_{g,j} = \mu_{g,j}$ and $\sigma_{g,j} \to 0$ not $\sigma_{g,j}^2 = \sigma_{g,j}^2$. Hence, we think Eq. (4.10) is not a proper similarity measure for two Gaussian distributions.

4.8 Summary

In this work, assuming that large scale data is modeled by Gaussian distributions, we studied the problem of similarity search over non-correlated Gaussian distributions based on KL-divergence. We analyzed the mathematical properties of KL-divergence of Gaussian distributions. Based on the analysis, we proposed two types of approaches to efficiently and effectively process top-$k$ similarity search over Gaussian distributions, which returns the $k$ most similar ones to a given query Gaussian distribution. They utilize the notions
of rank aggregation and skyline queries, respectively. We demonstrated the efficiency and effectiveness of our approaches through a comprehensive experimental performance study.
Chapter 5

Conclusions and Future Work

In this chapter, we conclude this thesis in Section 5.1 and present several interesting future directions in Section 5.2.

5.1 Conclusions

In this thesis, we studied uncertain data management due to the increasing uncertain data in a wide variety of fields in recent decades. Uncertainty can occur for different reasons and can be represented by a number of models. In particular, probabilistic model is utilized by many researchers to represent uncertainties using probability distributions. Among the three categories of probabilistic model, which are table-based, tuple-based, and attribute-based, we focused our attention on the third category, attribute-based, and further concentrate on the continuous case where the uncertain attribute is represented by a continuous probability density function, specifically, Gaussian distribution. In other words, we assumed that each uncertain object in the databases is described by a Gaussian distribution.

We considered three types of queries over probabilistic data with Gaussian distributions. The first one is probabilistic range query, which searches for objects within the given range with probabilities no less than a specified probability threshold. The query object
can be either a certain point or an uncertain object represented by a Gaussian distribution. In the second work, we investigated expected nearest neighbor search based on the expected distance. The query object considered in this work is a certain point. The result objects are ones that have the $k$ smallest expected distance to the query point. Finally, our third work studied similarity search over Gaussian distributions. The query object is also a Gaussian distribution. We use the KL-divergence as the similarity measure between two Gaussian distributions and return the top-$k$ similar Gaussian distributions as the result.

Answering queries over uncertain databases poses a number of challenges since managing uncertainty usually means costly probability computations. Hence, we developed efficient solutions for these three problems and conducted comprehensive performance studies using both synthetic and real datasets. We summarized each of our proposed solutions as follows.

5.1.1 Probabilistic Range Queries

In this work, we studied probabilistic range queries over uncertain data represented by Gaussian distributions. We assumed that the locations of data objects are represented by Gaussian distributions and the location of the query object is either fixed or follows a multi-dimensional Gaussian distribution. Under this setting, we defined two types of probabilistic range queries with respect to the query object and called them PRQ-$P$ and PRQ-$G$ queries, respectively.

To reduce expensive probability computations, we proposed a set of filtering techniques to avoid unnecessary computations. We designed a number of filtering polices for both PRQ-$P$ and PRQ-$G$ queries based on approximated regions that are derived from our analysis of properties of Gaussian distribution. The proposed filtering techniques are very effective in reducing the number of objects that need to be verified by probability computation and thus can save computation cost.

Furthermore, we proposed a novel indexing method called G-tree to accelerate query processing. We developed the indexing method by extending the existing R-tree and improved it based on our analysis of Gaussian distribution. The indexing method organizes
all the objects very effectively and enables filtering unpromising objects by groups rather than individually. Hence, the performance of query processing can be greatly enhanced.

The experimental results show that at least 99.5% of all the objects in the dataset can be pruned on average and over 58% of them are real result objects after evaluation by exact probability computation. This demonstrates the great pruning power of our proposed filtering techniques. Moreover, we equipped R-tree with our filtering techniques, called FR-tree, to evaluate the performance of our proposed indexing method G-tree. In the experiments, we compared them by varying the dataset size, the query range, the probability threshold, and the dimensionality. In all cases, G-tree outperforms FR-tree and shows great scalability and stability.

### 5.1.2 Expected Nearest Neighbor Search

In this work, we considered $k$-expected nearest neighbor search over objects represented by Gaussian distributions. This query finds the top-$k$ objects that are nearest to a given query point based on their expected distances to the query point. In other words, the result objects are ones that have the $k$ smallest expected distance to the query point. Since the naïve approach of performing sequential scan is computationally expensive for solving this problem, we proposed novel solutions to support efficient query processing.

We analyzed properties of expected distance on Gaussian distribution mathematically and derived the lower bound and upper bound of the distance. Based on our analysis, we proposed three novel approaches, AVG, PLB, and PLUB, to efficiently prune unpromising objects without computing their actual expected distances. We only compute actual expected distances for candidate objects and finally return the top-$k$ smallest ones. To further improve the performance, we utilized R-tree to index objects and their lower bound distances and upper bound distances.

Both the lower bound and upper bound are described by the distance between the average point of a Gaussian distribution and the query point plus a non-negative minimum or maximum value that depends on the covariance matrix of the Gaussian distribution. The first one, AVG uses simply the distance between the average point and the query point as
the lower bound to prune unpromising objects whose lower bound distances are greater than the actual expected distances of top-\(k\) candidates. This approach is rather simple, but it enables us to pay less cost on storing information about the covariance matrix in the R-tree. The second approach, PLB employs the more precise lower bound for filtering and needs to maintain more information about the covariance matrix. Finally, the third one, PLUB exploits both the lower bound and upper bound to prune non-candidates as much as possible. The non-candidates here are objects whose lower bound distances are greater than upper bound distances of candidate objects. Correspondingly, PLUB stores the most information about the covariance matrix. In other words, they are tradeoffs in different degrees between computation cost and storage cost.

In the experiments, we compared the performance of our proposed approaches AVG, PLB, and PLUB, by varying the three parameters, \(k\), dataset size, and dimensionality. All the approaches show good scalability and stability over these parameters. We found that among the three approaches, PLUB achieves the best efficiency while AVG is better than PLB if \(k\) is large, and PLB is a better choice in the case of a large data size.

5.1.3 Similarity Search

In this work, we investigated similarity search on uncertain data modeled in non-correlated Gaussian distributions, where there are no correlations between dimensions and their covariance matrices are diagonal. The query object is also represented by a non-correlated Gaussian distribution. We employed KL-divergence to measure the similarity between two Gaussian distributions. This query returns the top-\(k\) Gaussian distributions that are similar to a given query Gaussian distribution based on KL-divergence. In other words, the result objects are Gaussian distributions that have the top-\(k\) smallest KL-divergence with the query Gaussian distribution.

We analyzed mathematically KL-divergence of Gaussian distributions and derived its divisionally monotonous properties. Based on the analysis, we proposed two types of approaches, TA-based and Skyline-based, to efficiently and effectively solve this problem. They utilize the notions of rank aggregation and skyline queries, respectively. The first
type presorts all objects in the database on their attributes and computes result objects by merging candidates from each presorted list. We extended the existing TA algorithm and developed two novel algorithms, CTA and PTA. The second one transforms the problem to the computation of dynamic skyline queries. We extended and modified the BBS algorithm and developed a novel algorithm called SKY to solve this problem.

In our experimental study, we evaluated the effects of $k$, dataset size, dimensionality, and data distribution on our proposed three approaches, CTA, PTA, and SKY. The experimental results show that all of our approaches performs better than the naïve approach Scan. Among them, SKY demonstrates the best and the performance of PTA and CTA is between SKY and Scan, with that CTA performs slightly better. Moreover, we found that our proposed approaches show great stability over different data distributions.

5.2 Future Work

In this thesis, we considered the attribute-level uncertainty represented by Gaussian distribution and our proposed approaches are mainly based the spatial index R-tree. As one future direction, we can consider other uncertainty models such as other probability distributions and the discrete uncertainty model. Furthermore, we can consider the Gaussian Mixture model, which can be used to represent arbitrary complex probability distributions. Another future direction is to develop novel index structures that are able to be adapted to multiple uncertainty models. For instance, we can combine the power of R-tree with that of other indexing methods such as Grid-based index structure and Quadtree. In addition, we plan to design an integrated query framework that can support multiple types of queries at the same time.

Other interesting research directions include applying other types of queries such as skyline queries, join queries, reverse queries, and aggregate queries, over probabilistic data with Gaussian distributions and defining novel types of queries by observing and pondering over emerging real-world applications. Moreover, we can consider queries over probabilistic data with Gaussian distributions in distributed environments and road networks.
In the following sections, we present specific future work for each of our three work.

5.2.1 Probabilistic Range Queries

In the current implementation, the node split policy of G-tree follows that of R*-tree and the computation of the four penalty metrics (area, margin, overlap and centroid distance) used for splitting is based on that of TPR-tree. As the future work, the structure of G-tree can be optimized for more efficient query processing by taking advantage of its features. Furthermore, we plan to develop novel indexing methods based on other index structures such as the Grid-based index structure.

5.2.2 Expected Nearest Neighbor Search

Currently, we use directly R-tree to index Gaussian distributions and their lower bound and upper bound distances. In the future, we consider improving and extending R-tree for expected nearest neighbor search over Gaussian distributions. We also plan to consider other indexing methods such as the Grid-based index structure. Moreover, we consider adapting the indexing method proposed in our first work to support query processing in this work. In addition, we plan to extend our work to distance measures other than the expected distance.

5.2.3 Similarity Search

In the future, we plan to improve index structures to reduce preprocessing cost. In this work, we use directly R-tree to index Gaussian distributions. Therefore, in the future, we consider improving and extending R-tree for similarity search over Gaussian distributions. We also plan to consider other indexing methods such as the Grid-based index structure besides R-tree. Furthermore, we will study the similarity search problem in the general case of multi-dimensional Gaussian distributions and use similarity measures other than KL-divergence.
Appendix A

Appendix for Chapter 2

A.1 Proof of Lemma 2.3

As shown in Fig. A.1(a), consider the case where the minimum distance between $bb_i(\rho)$ and $bb_q(\rho)$ is $\delta$. The space outside the $\rho$-region of $o_i$ (resp. $q$) is divided into two parts by the line $l_a$ (resp. $l_d$) equally, both with an existence probability of $(1 - \rho)/2$. Assuming the probability that $o_i$ (resp. $q$) is located in the dashed area between $l_a$ and $l_b$ (resp. $l_c$ and $l_d$) excluding the half part of $\rho$-region is $\alpha$ (resp. $\beta$), the probability that $o_i$ lies within the left part of $l_b$ and $q$ lies within the left part of $l_c$ is $((1 - \rho)/2 + \rho + \alpha)((1 - \rho)/2 - \beta)$. When $o_i$ lies within the right part of $l_b$ ($q$ can be located both in the right and left part of $l_c$), the probability is $((1 - \rho)/2 - \alpha)$. Thus, the maximal probability of $\|x_o - x_q\| \leq \delta$ can be calculated by summing up the two probabilities, resulting in

$$
Pr(\|x_o - x_q\| \leq \delta) < ((1 - \rho)/2 + \rho + \alpha)((1 - \rho)/2 - \beta) + ((1 - \rho)/2 - \alpha)
$$

$$
= (3 - 2\rho - \rho^2)/4 - (\alpha + \beta)(1 + \rho)/2 - \alpha\beta
$$

$$
< (3 - 2\rho - \rho^2)/4.
$$

□
Appendix A. Appendix for Chapter 2

Figure A.1: Proof of maximal probability for PRQ-G

A.2 Proof of Lemma 2.4

Assume that the distance between the two mean locations is exactly $\delta$ as illustrated in Fig. A.1(b). $o_i$ (resp. $q$) has a probability of 0.5 to be located in both left and right part of the line $l_o$ (resp. $l_q$). $\|x_o - x_q\| \leq \delta$ happens in three cases: (1) Both $o_i$ and $q$ distribute in the left part of $l_o$ and $l_q$. (2) Both $o_i$ and $q$ distribute in the right part of $l_o$ and $l_q$. (3) $o_i$ distributes in the right part of $l_o$ and $q$ distributes in the left part of $l_q$. Each case has a probability of $0.5 \times 0.5$. Hence, the maximal probability of $\|x_o - x_q\| \leq \delta$ is $0.5 \times 0.5 \times 3 = 0.75$. \qed
Appendix B

Appendix for Chapter 3

B.1 The Proof of Lower bound and upper bound of $\Delta^2$

Since $\Delta^2 = \int \|x - \mu\|^2 \cdot p(x) dx$ and $p^+(x) \leq p(x) \leq p^\top(x)$, we have

$$\Delta^2_{\text{min}} = \int \|x - \mu\|^2 \cdot p^+(x) dx \leq \Delta^2 \leq \int \|x - \mu\|^2 \cdot p^\top(x) dx = \Delta^2_{\text{max}}.$$ 

Here,

$$p^+(x) = \frac{1}{(2\pi)^{d/2}|\Sigma|^{1/2}} \exp \left[-\frac{\|x\|^2}{2\lambda^\perp}\right]$$

$$p^\top(x) = \frac{1}{(2\pi)^{d/2}|\Sigma|^{1/2}} \exp \left[-\frac{\|x\|^2}{2\lambda^\top}\right]$$

where $\lambda^\perp$ and $\lambda^\top$ are the minimum and maximum eigenvalues of $\Sigma$, respectively. We first derive $\Delta^2_{\text{min}}$ and then apply the same process to $\Delta^2_{\text{max}}$.

Assume that in a $d$-dimensional space a vector $x$ is represented by $(x_1, \ldots, x_d)$. By assuming $y = x - \mu$, we have

$$\Delta^2_{\text{min}} = \int \|x - \mu\|^2 \cdot \frac{1}{(2\pi)^{d/2}|\Sigma|^{1/2}} \exp \left[-\frac{\|x - \mu\|^2}{2\lambda^\perp}\right] dx$$

$$= \frac{1}{(2\pi)^{d/2}|\Sigma|^{1/2}} \int \|y\|^2 \cdot \exp \left[-\frac{\|y\|^2}{2\lambda^\perp}\right] dy. \quad (B.1)$$
Appendix B: Appendix for Chapter 3

According to [113], we transform Cartesian coordinates to spherical polar coordinates by

\[
\begin{align*}
y_1 &= r \cos \theta_1 \ldots \cos \theta_{d-2} \cos \theta_{d-1} = rc_1 \ldots c_{d-2}c_{d-1} \\
y_2 &= r \cos \theta_1 \ldots \cos \theta_{d-2} \sin \theta_{d-1} = rc_1 \ldots c_{d-2}s_{d-1} \\
\vdots \\
y_k &= r \cos \theta_1 \ldots \cos \theta_{d-k} \sin \theta_{d-k+1} = rc_1 \ldots c_{d-k}s_{d-k+1} \\
\vdots \\
y_1 &= r \sin \theta_1 = rs_1.
\end{align*}
\]

The Jacobian is \((-1)^{d-1}c_1^{d-2}c_2^{d-3} \ldots c_{d-2} \). Then we have

\[
\int ||y||^2 \cdot \exp\left[\frac{-||y||^2}{2 \lambda^2}\right] dy = \int_{-\infty}^{+\infty} \ldots \int_{-\infty}^{+\infty} (\sum_{i=1}^{d} y_i^2) \cdot \exp\left[\frac{-1}{2 \lambda^2} \sum_{i=1}^{d} y_i^2\right] dy_1 \ldots dy_d \\
= \int_{0}^{+\infty} \int_{-\pi/2}^{+\pi/2} \int_{-\pi/2}^{+\pi/2} r^2 \cdot \exp\left[\frac{-r^2}{2 \lambda^2}\right] \cdot \|(-1)^{d-1}c_1^{d-2}c_2^{d-3} \ldots c_{d-2}\| drd\theta_1 \ldots d\theta_{d-1} \\
= \int_{0}^{r^{d+1}} \exp\left[\frac{-r^2}{2 \lambda^2}\right] dr \cdot \int_{-\pi/2}^{+\pi/2} c_1^{d-2}d\theta_1 \ldots \int_{-\pi/2}^{+\pi/2} c_{d-2}d\theta_{d-2} \cdot \int_{0}^{2\pi} r^2 d\theta_{d-1}. \quad (B.2)
\]

According to [114],

\[
\int_{0}^{\infty} x^n e^{-\alpha x} dx = \frac{b}{\alpha} \left(\frac{n+1}{b}\right).
\]

By letting \(n = d + 1, a = \frac{1}{2 \lambda^2}\), and \(b = 2\), we can obtain

\[
\int_{0}^{\infty} r^{d+1} \cdot \exp\left[\frac{-r^2}{2 \lambda^2}\right] dr = \frac{1}{2} (2 \lambda^2 \alpha)^{d+2} \gamma\left(\frac{d+2}{2}\right).
\]

Then Eq. (B.2) becomes

\[
\int ||y||^2 \cdot \exp\left[\frac{-||y||^2}{2 \lambda^2}\right] dy = \frac{1}{2} (2 \lambda^2 \alpha)^{d+2} \gamma\left(\frac{d+2}{2}\right) \cdot \int_{\pi/2}^{+\pi/2} c_1^{d-2}d\theta_1 \ldots \int_{\pi/2}^{+\pi/2} c_{d-2}d\theta_{d-2} \cdot \int_{0}^{2\pi} r^2 d\theta_{d-1} \\
= (2 \lambda^2 \alpha)^{d+2} \gamma\left(\frac{d+2}{2}\right) \cdot \int_{\pi/2}^{+\pi/2} c_1^{d-2}d\theta_1 \ldots \int_{\pi/2}^{+\pi/2} c_{d-2}d\theta_{d-2} \cdot \pi. \quad (B.3)
\]
Based on properties of Gamma function, the following statements hold.

1. If \( d \) is odd, then
   \[
   \gamma\left(\frac{d + 2}{2}\right) = \frac{d}{2} \cdot \frac{d - 2}{2} \cdots \frac{1}{2} \cdot \sqrt{\pi}. \tag{B.4}
   \]

2. If \( d \) is even, then
   \[
   \gamma\left(\frac{d + 2}{2}\right) = \frac{d}{2} \cdot \frac{d - 2}{2} \cdots \frac{2}{2} \cdot 1. \tag{B.5}
   \]

Also, when \( n > 0 \) we have
\[
\int_{-\pi/2}^{+\pi/2} \cos^n \theta d\theta = \left[ \frac{\cos^{n-1} \theta \sin \theta}{n} \right]_{-\pi/2}^{\pi/2} + \frac{n-1}{n} \int_{-\pi/2}^{\pi/2} \cos^{n-2} \theta d\theta
= \frac{n-1}{n} \int_{-\pi/2}^{\pi/2} \cos^{n-2} \theta d\theta.
\]

By deduction, the following statements hold.

1. If \( d \) is odd, then
   \[
   \int_{-\pi/2}^{+\pi/2} \cos^n \theta d\theta = \frac{n-1}{n} \cdot \frac{n-3}{n-2} \cdots \frac{4}{3} \cdot \frac{2}{3} \cdot 2.
   \]

2. If \( d \) is even, then
   \[
   \int_{-\pi/2}^{+\pi/2} \cos^n \theta d\theta = \frac{n-1}{n} \cdot \frac{n-3}{n-2} \cdots \frac{3}{4} \cdot \frac{1}{2} \cdot \pi.
   \]

Thus, we have

1. If \( d \) is odd, then
   \[
   \int_{-\pi/2}^{+\pi/2} c_{d-2}^{d-2} \theta_1 \cdots \int_{-\pi/2}^{+\pi/2} c_{d-2}^{d-2} d\theta_{d-2}
   = \left( \frac{d-3}{d-2} \cdot \frac{d-5}{d-4} \cdots \frac{2}{3} \cdot \frac{2}{3} \cdot 2 \right) \cdot \left( \frac{d-4}{d-3} \cdot \frac{d-6}{d-5} \cdots \frac{3}{4} \cdot \frac{1}{2} \cdot \pi \right) \cdots 2
   = \frac{2\pi}{d-2} \cdots \frac{2\pi}{3} \cdot 2. \tag{B.6}
   \]
2. If $d$ is even, then

\[
\int_{-\pi/2}^{+\pi/2} c_{d-2} d\theta_1 \cdots \int_{-\pi/2}^{+\pi/2} c_{d-2} d\theta_{d-2} = \left( \frac{d-3}{d-2} \cdot \frac{d-5}{d-4} \cdots \frac{3}{2} \cdot \pi \right) \cdot \left( \frac{d-4}{d-3} \cdot \frac{d-6}{d-5} \cdots \frac{2}{3} \cdot \frac{1}{2} \right) \cdot \pi \cdot 2 = \frac{2\pi}{d-2} \cdot \frac{3\pi}{4} \cdot \pi. \tag{B.7}
\]

Based on Eq. (B.4), Eq. (B.5), Eq. (B.6), and Eq. (B.7), we have

1. If $d$ is odd, then

\[
\gamma \left( \frac{d+2}{2} \right) \cdot \int_{-\pi/2}^{+\pi/2} c_{1}^{d-2} d\theta_1 \cdots \int_{-\pi/2}^{+\pi/2} c_{d-2} d\theta_{d-2} = \left( \frac{d}{2} \cdot \frac{d-2}{2} \cdots \frac{1}{2} \cdot \sqrt{\pi} \right) \cdot \left( \frac{2\pi}{d-2} \cdot \frac{2\pi}{3} \cdot \frac{2}{2} \right) = \frac{d}{2} \pi^{d-1}. \tag{B.8}
\]

2. If $d$ is even, then

\[
\gamma \left( \frac{d+2}{2} \right) \cdot \int_{-\pi/2}^{+\pi/2} c_{1}^{d-2} d\theta_1 \cdots \int_{-\pi/2}^{+\pi/2} c_{d-2} d\theta_{d-2} = \left( \frac{d}{2} \cdot \frac{d-2}{2} \cdots \frac{2}{2} \cdot 1 \right) \cdot \left( \frac{2\pi}{d-2} \cdot \frac{2\pi}{4} \cdot \pi \right) = \frac{d}{2} \pi^{d-1}. \tag{B.8}
\]

In other words, the following equation holds no matter $d$ is odd or even.

\[
\gamma \left( \frac{d+2}{2} \right) \cdot \int_{-\pi/2}^{+\pi/2} c_{1}^{d-2} d\theta_1 \cdots \int_{-\pi/2}^{+\pi/2} c_{d-2} d\theta_{d-2} = \frac{d}{2} \pi^{d-1}. \tag{B.8}
\]

Then Eq. (B.3) becomes

\[
\int \|y\|^2 \cdot \exp \left[ -\frac{\|y\|^2}{2\lambda^2} \right] \, dy = (2\lambda \lambda^b)^{d-3/2} \cdot \frac{d}{2} \pi^{d-1} = d \cdot (\lambda \lambda^b)^{d+1} \cdot (2\pi)^{d}. \tag{B.8}
\]

Based on Eq. (B.1), and Eq. (B.8), we have

\[
\Delta_{\text{min}}^2 = \frac{1}{(2\pi)^d/|\Sigma|^{1/2}} \cdot d \cdot (\lambda \lambda^b)^{d+1} \cdot (2\pi)^{d} = \frac{d \cdot (\lambda \lambda^b)^{1+d/2}}{|\Sigma|^{1/2}}. 
\]
Similarly, we can prove

$$\Delta^2_{\text{max}} = \frac{d \cdot (\lambda' o p)^{1+d/2}}{|\Sigma|^{1/2}}.$$ 

Hence, we have proved the lower bound and upper bound of $\Delta^2$. □
Appendix C

Appendix for Chapter 4

C.1 Proof of Lemma 4.1

C.1.1 Case 1: $\mathcal{D}_{KL}(p||q)$

Assume $\sigma_{p,j}^2 - \sigma_{q,j}^2 = C_1$ and $\sigma_{q,j}^2 - \sigma_{p',j}^2 = C_2$, i.e. $\sigma_{p,j}^2 = \sigma_{q,j}^2 + C_1$ and $\sigma_{p',j}^2 = \sigma_{q,j}^2 - C_2$. Then $0 < C_1 \leq C_2 < \sigma_{q,j}^2$.

Let $\Delta_1 = \mathcal{D}_{KL}(p||q) - \mathcal{D}_{KL}(p'||q)$. Then

$$\Delta_1 = \frac{1}{2} \ln \frac{\sigma_{q,j}^2 - C_2}{\sigma_{q,j}^2 + C_1} + \frac{C_1 + C_2}{2\sigma_{q,j}^2},$$

$$\frac{\partial \Delta_1}{\partial \sigma_{q,j}^2} = \frac{(C_1 + C_2)[(C_2 - C_1)\sigma_{q,j}^2 + C_1C_2]}{2\sigma_{q,j}^4(\sigma_{q,j}^2 - C_2)(\sigma_{q,j}^2 + C_1)} > 0.$$

Since $\Delta_1|_{\sigma_{q,j}^2\to\infty} = 0$, $\Delta_1 < 0$ holds for all $\sigma_{q,j}^2$, i.e., $\mathcal{D}_{KL}(p||q) < \mathcal{D}_{KL}(p'||q)$.

C.1.2 Case 2: $\mathcal{D}_{KL}(q||p)$

Since $|\mu_{p,j} - \mu_{q,j}| = |\mu_{p',j} - \mu_{q,j}|$, we use $|\mu_{p,j} - \mu_{q,j}|$ to represent both of them. Assume $\sigma_{p,j}^2 - \sigma_{q,j}^2 - (\mu_{p,j} - \mu_{q,j})^2 = C_1$ and $\sigma_{q,j}^2 - \sigma_{p',j}^2 - (\mu_{p,j} - \mu_{q,j})^2 = C_2$, i.e., $\sigma_{p,j}^2 = \sigma_{q,j}^2 + (\mu_{p,j} - \mu_{q,j})^2 + C_1$ and $\sigma_{p',j}^2 = \sigma_{q,j}^2 + (\mu_{p,j} - \mu_{q,j})^2 - C_2$. Then $0 < C_1 \leq C_2 < \sigma_{q,j}^2 + (\mu_{p,j} - \mu_{q,j})^2$. 96
Let $\Delta_2 = D_{KL}(q||p) - D_{KL}(q||p')$ and $\alpha = \sigma_{q,j}^2 + (\mu_{p,j} - \mu_{q,j})^2$. Then

$$\Delta_2 = \frac{1}{2} \ln \frac{\alpha + C_1}{\alpha - C_2} - \frac{\alpha(C_1 + C_2)}{2(\alpha + C_1)(\alpha - C_2)},$$

$$\frac{\partial \Delta_2}{\partial \sigma_{q,j}^2} = \frac{(C_1 + C_2)[\alpha(C_2 - C_1) + 2C_1C_2]}{2(\alpha - C_2)^2(\alpha + C_1)^2} > 0.$$

Since $\Delta_2|_{\sigma_{q,j}^2 \to 0} = 0$, $\Delta_2 < 0$ holds for all $\sigma_{q,j}^2$, i.e., $D_{KL}(q||p) < D_{KL}(q||p')$.

C.2 Proof of Lemma 4.4

Let the mmdist of $g$ be mmdist$_g$. Due to the definition of mmdist, for any equal-KLD point $g'$ of $g$, mmdist$_g \geq |\mu_{g'} - \mu_g| + |\sigma_{g'}^2 - \sigma_q^2|$ holds. Since mindist$_e$, the mindist of an entry $e$, satisfies mindist$_e = |\mu_e - \mu_g| + |\sigma_e^2 - \sigma_q^2| > mmdist_g$, we have

$$|\mu_e - \mu_g| + |\sigma_e^2 - \sigma_q^2| > |\mu_{g'} - \mu_g| + |\sigma_{g'}^2 - \sigma_q^2|. \quad (C.1)$$

Given $q$ and the KLD of $g$ and $q$, and assuming

$$D_{KL}^1(g||q) = \frac{1}{2} \left[ \frac{(\mu_g - \mu_q)^2}{\sigma_q^2} + \frac{\sigma_q^2}{\sigma_g^2} - \ln \frac{\sigma_q^2}{\sigma_g^2} - 1 \right] = C,$$

when $\sigma_q^2 = \sigma_g^2$, $(\mu_g - \mu_q)^2$ takes the maximum $2C\sigma_q^2$.

The reason is as follows. We can easily find that the function $f(x) = x - \ln x$ takes the minimum when $x = 1$. Therefore, $\frac{\sigma_g^2}{\sigma_q^2} - \ln \frac{\sigma_g^2}{\sigma_q^2}$ takes the minimum when $\frac{\sigma_g^2}{\sigma_q^2} = 1$. In other words, when $\sigma_q^2 = \sigma_g^2$, $(\mu_g - \mu_q)^2$ is the maximum. We prove the lemma in the following two cases.

1. $(\mu_e - \mu_g)^2 > 2C\sigma_q^2$:

Consider $g'$, the equal-KLD point $g'$ satisfying $|\mu_{g'} - \mu_g| = \sqrt{2C}\sigma_q$, and $\sigma_{g'}^2 = \sigma_q^2$. Since $|\mu_e - \mu_g| > \sqrt{2C}\sigma_q = |\mu_{g'} - \mu_g|$, and $|\sigma_e^2 - \sigma_q^2| \geq 0 = |\sigma_{g'}^2 - \sigma_q^2|$, it is obvious that $e$ is dynamically dominated by $g'$, i.e., $e$ is dynamically KLD-dominated by $g$. 


2. Otherwise:

Consider $g$’s equal-KLD point $g'$ satisfying $\mu_{g'} = \mu_e$. According to eq.(C.1), we have $|\sigma^2_e - \sigma^2_q| > |\sigma^2_{g'} - \sigma^2_q|$. In other words, $|\mu_e - \mu_q| = |\mu_{g'} - \mu_q|$, and $|\sigma^2_e - \sigma^2_q| > |\sigma^2_{g'} - \sigma^2_q|$. Therefore, $e$ is dynamically dominated by $g'$, i.e., $e$ is dynamically KLD-dominated by $g$. 

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