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Ranking Spatial Data by Quality Preferences

Man Lung Yiu, Hua Lu, Member, IEEE, Nikos Mamoulis, and Michail Vaitis

Abstract—A spatial preference query ranks objects based on the qualities of features in their spatial neighborhood. For example, using a real estate agency database of flats for lease, a customer may want to rank the flats with respect to the appropriateness of their location, defined after aggregating the qualities of other features (e.g., restaurants, cafes, hospital, market, etc.) within their spatial neighborhood. Such a neighborhood concept can be specified by the user via different functions. It can be an explicit circular region within a given distance from the flat. Another intuitive definition is to assign higher weights to the features based on their proximity to the flat. In this paper, we formally define spatial preference queries and propose appropriate indexing techniques and search algorithms for them. Extensive evaluation of our methods on both real and synthetic data reveals that an optimized branch-and-bound solution is efficient and robust with respect to different parameters.

Index Terms—Query processing, spatial databases.

1 INTRODUCTION

Spatial database systems manage large collections of geographic entities, which apart from spatial attributes contain nonspatial information (e.g., name, size, type, price, etc.). In this paper, we study an interesting type of preference queries, which select the best spatial location with respect to the quality of facilities in its spatial neighborhood.

Given a set \( D \) of interesting objects (e.g., candidate locations), a top-\( k \) spatial preference query retrieves the \( k \) objects in \( D \) with the highest scores. The score of an object is defined by the quality of features (e.g., facilities or services) in its spatial neighborhood. As a motivating example, consider a real estate agency office that holds a database with available flats for lease. Here “feature” refers to a class of objects in a spatial map such as specific facilities or services. A customer may want to rank the contents of this database with respect to the quality of their locations, quantified by aggregating nonspatial characteristics of other features (e.g., restaurants, cafes, hospital, market, etc.) in the spatial neighborhood of the flat (defined by a spatial range around it). Quality may be subjective and query-parametric. For example, a user may define quality with respect to nonspatial attributes of restaurants around it (e.g., whether they serve seafood, price range, etc.).

As another example, the user (e.g., a tourist) wishes to find a hotel \( p \) that is close to a high-quality restaurant and a high-quality cafe. Fig. 1a illustrates the locations of an object data set \( D \) (hotels) in white, and two feature data sets: the set \( F_1 \) (restaurants) in gray, and the set \( F_2 \) (cafes) in black. Feature points are labeled by quality values that can be obtained from rating providers (e.g., http://www.zagat.com/). For the ease of discussion, the qualities are normalized to values in \([0, 1]\).

The score \( \tau(p) \) of a hotel \( p \) is defined in terms of: 1) the maximum quality for each feature in the neighborhood region of \( p \), and 2) the aggregation of those qualities.

A simple score instance, called the range score, binds the neighborhood region to a circular region at \( p \) with radius \( \epsilon \) (shown as a circle), and the aggregate function to SUM. For instance, the maximum quality of gray and black points within the circle of \( p_1 \) are 0.9 and 0.6, respectively, so the score of \( p_1 \) is \( \tau(p_1) = 0.9 + 0.6 = 1.5 \). Similarly, we obtain \( \tau(p_2) = 1.0 + 0.1 = 1.1 \) and \( \tau(p_3) = 0.7 + 0.7 = 1.4 \). Hence, the hotel \( p_1 \) is returned as the top result.

In fact, the semantics of the aggregate function is relevant to the user’s query. The SUM function attempts to balance the overall qualities of all features. For the MAX function, the top result becomes \( p_2 \), with the score \( \tau(p_2) = \max(0.7, 0.7) = 0.7 \). It ensures that the top result has reasonably high qualities in all features. For the MIN function, the top result is \( p_2 \), with \( \tau(p_2) = \min(1.0, 0.1) = 0.1 \). It is used to optimize the quality in a particular feature, but not necessarily all of them.

The neighborhood region in the above spatial preference query can also be defined by other score functions. A meaningful score function is the influence score (see Section 4). As opposed to the crisp radius \( \epsilon \) constraint in the range score, the influence score smoothens the effect of \( \epsilon \) and assigns higher weights to cafes that are closer to the hotel.

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Object ranking is a popular retrieval task in various applications. In relational databases, we rank tuples using an aggregate score function on their attribute values [2]. For example, a real estate agency maintains a database that contains information of flats available for rent. A potential customer wishes to view the top 10 flats with the largest sizes and lowest prices. In this case, the score of each flat is expressed by the sum of two qualities: size and price, after normalization to the domain $[0, 1]$ (e.g., 1 means the largest size and the lowest price). In spatial databases, ranking is often associated to nearest neighbor (NN) retrieval. Given a query location, we are interested in retrieving the set of nearest objects to it that qualify a condition (e.g., restaurants). Assuming that the set of interesting objects is indexed by an R-tree [3], we can apply distance bounds and traverse the index in a branch-and-bound fashion to obtain the answer [4].

Nevertheless, it is not always possible to use multi-dimensional indexes for top-$k$ retrieval. First, such indexes break down in high-dimensional spaces [5], [6]. Second, top-$k$ queries may involve an arbitrary set of user-specified attributes (e.g., size and price) from possible ones (e.g., size, price, distance to the beach, number of bedrooms, floor, etc.) and indexes may not be available for all possible attribute combinations (i.e., they are too expensive to create and maintain). Third, information for different rankings to be combined (i.e., for different attributes) could appear in different databases (in a distributed database scenario) and unified indexes may not exist for them. Solutions for top-$k$ queries [7], [2], [8], [9] focus on the efficient merging of object rankings that may arrive from different (distributed) sources. Their motivation is to minimize the number of accesses to the input rankings until the objects with the top-$k$ aggregate scores have been identified. To achieve this, upper and lower bounds for the objects seen so far are maintained while scanning the sorted lists.

In the following sections, we first review the R-tree, which is the most popular spatial access method and the NN search algorithm of [4]. Then, we survey recent research of feature-based spatial queries.

### 2.1 Spatial Query Evaluation on R-Trees

The most popular spatial access method is the R-tree [3], which indexes minimum bounding rectangles (MBRs) of objects. Fig. 2 shows a set $D = \{p_1, \ldots, p_8\}$ of spatial objects (e.g., points) and an R-tree that indexes them. R-trees can efficiently process main spatial query types, including spatial range queries, nearest neighbor queries, and spatial joins. Given a spatial region $W$, a spatial range query retrieves from $D$ the objects that intersect $W$. For instance, consider a range query that asks for all objects within the shaded area in Fig. 2. Starting from the root of the tree, the query is processed by recursively following entries, having MBRs intersect the query region, thus the subtree pointed by $e_1$.

![Fig. 2. Spatial queries on R-trees.](image-url)
cannot contain any query result. In contrast, $e_2$ is followed by the algorithm and the points in the corresponding node are examined recursively to find the query result $p_7$.

A nearest neighbor query takes as input a query object $q$ and returns the closest object in $D$ to $q$. For instance, the nearest neighbor of $q$ in Fig. 2 is $p_7$. Its generalization is the $k$-NN query, which returns the $k$ closest objects to $q$, given a positive integer $k$. NN (and $k$-NN) queries can be efficiently processed using the best-first (BF) algorithm of [4], provided that $D$ is indexed by an R-tree. A min-heap $H$ which organizes R-tree entries based on the (minimum) distance of each leaf entry to the root entries. In order to find the NN of $q$ in Fig. 2, BF first inserts to $H$ entries $e_1$, $e_2$, $e_3$, and their distances to $q$. Then, the nearest entry $e_2$ is retrieved from $H$ and objects $p_1$, $p_7$, $p_8$ are inserted to $H$. The next nearest entry in $H$ is $p_7$, which is the nearest neighbor of $q$. In terms of I/O, the BF algorithm is shown to be no worse than any NN algorithm on the same R-tree [4].

The aggregate R-tree (aR-tree) [10] is a variant of the R-tree, where each nonleaf entry augments an aggregate measure for some attribute value (measure) of all points in its subtree. As an example, the tree shown in Fig. 2 can be upgraded to a MAX aR-tree over the point set, if entries $e_1, e_2, e_3$ contain the maximum measure values of sets $\{p_2, p_3\}, \{p_1, p_7, p_8\}, \{p_4, p_5, p_6\}$, respectively. Assume that the measure values of $p_1, p_7, p_8$ are 0.2, 0.1, 0.4, respectively. In this case, the aggregate measure augmented in $e_3$ would be $\max\{0.2, 0.1, 0.4\} = 0.4$. In this paper, we employ MAX aR-trees for indexing the feature data sets (e.g., restaurants), in order to accelerate the processing of top-$k$ spatial preference queries.

Given a feature data set $F$ and a multidimensional region $R$, the range top-$k$ query selects the tuples (from $F$) within the region $R$ and returns only those with the $k$ highest qualities. Hong et al. [11] indexed the data set by a MAX aR-tree and developed an efficient tree traversal algorithm to answer the query. Instead of finding the best $k$ qualities from $F$ in a specified region, our (range score) query considers multiple spatial regions based on the points from the object data set $D$, and attempts to find out the best $k$ regions (based on scores derived from multiple feature data sets $F_c$).

### 2.2 Feature-Based Spatial Queries

Xia et al. [12] solved the problem of finding top-$k$ sites (e.g., residential buildings). As an example, Fig. 3a shows a set of sites (white points), a set of features (black points with weights), such that each line links a feature point to its nearest site. The influence of a site $p_i$ is defined by the sum of weights of feature points having $p_i$ as their closest site. For instance, the score of $p_1$ is $0.9 + 0.5 = 1.4$. Similarly, the scores of $p_2$ and $p_1$ are 1.5 and 1.2, respectively. Hence, $p_2$ is returned as the top-1 influential site.

Related to top-$k$ influential sites query are the optimal location queries studied in [13], [14]. The goal is to find the location in space (not chosen from a specific set of sites) that minimizes an objective function. In Figs. 3b and 3c, feature points and existing sites are shown as black and gray points, respectively. Assume that all feature points have the same quality. The maximum influence optimal location query [13] finds the location (to insert the existing set of sites) with the maximum influence (as defined in [12]), whereas the minimum distance optimal location query [14] searches for the location that minimizes the average distance from each feature point to its nearest site. The optimal locations for both queries are marked as white points in Figs. 3b and 3c, respectively.

The techniques proposed in [12], [13], [14] are specific to the particular query types described above and cannot be extended for our top-$k$ spatial preference queries. Also, they deal with a single-feature data set whereas our queries consider multiple feature data sets.

Recently, novel spatial queries and joins [15], [16], [17], [18] have been proposed for various spatial decision support problems. However, they do not utilize nonspatial qualities of facilities to define the score of a location. Finally, [19], [20] studied the evaluation of textual location-based queries on spatial objects.

### 3 Spatial Preference Queries

Section 3.1 formally defines the top-$k$ spatial preference query problem and describes the index structures for the data sets. Section 3.2 studies two baseline algorithms for processing the query. Section 3.3 presents an efficient branch-and-bound algorithm for the query, and its further optimization is proposed in Section 3.4. Section 3.5 develops a specialized spatial join algorithm for evaluating the query. Finally, Section 3.6 extends the above algorithms for answering top-$k$ spatial preference queries involving other aggregate functions.

#### 3.1 Definitions and Index Structures

Let $F_c$ be a feature data set, in which each feature object $s \in F_c$ is associated with a quality $\omega(s)$ and a spatial point. We assume that the domain of $\omega(s)$ is the interval $[0, 1]$. As an example, the quality $\omega(s)$ of a restaurant $s$ can be obtained from a ratings provider.

Let $D$ be an object data set, where each object $p \in D$ is a spatial point. In other words, $D$ is the set of interesting points (e.g., hotel locations) considered by the user.

Given an object data set $D$ and $m$ feature data sets $F_1, F_2, \ldots, F_m$, the top-$k$ spatial preference query retrieves the $k$ points in $D$ with the highest score. Here, the score of an object point $p \in D$ is defined as

$$t^p = \text{ARG} \{ t^p_c | c \in [1, m] \}$$

\[ \text{ARG} \{ t^p_c | c \in [1, m] \} \]
where AGG is an aggregate function and \( \tau_\theta(p) \) is the (cth) component score of p with respect to the neighborhood condition \( \theta \) and the (cth) feature data set \( F_c \).

We proceed to elaborate the aggregate function and the component score function. Typical examples of the aggregate function AGG are: SUM, MIN, and MAX. We first focus on the case where AGG is SUM. In Section 3.6, we will discuss the generic scenario where AGG is an arbitrary monotone aggregate function.

An intuitive choice for the component score function \( \tau_\theta(p) \) is: the range score \( \tau^{\text{range}}_\theta(p) \), taken as the maximum quality \( \omega(s) \) of points \( s \in F_c \) that are within a given parameter distance \( \epsilon \) from \( p \), or 0 if no such point exists.

\[
\tau^{\text{range}}_\theta(p) = \max\{\omega(s) | s \in F_c \land \text{dist}(p, s) \leq \epsilon\} \cup \{0\}.
\] (2)

In our problem setting, the user requires that an object \( p \in D \) must not be considered as a result if there exists some \( F_c \) such that the neighborhood region of \( p \) does not contain any feature point of \( F_c \).

There are other choices for the component score function \( \tau_\theta(p) \). One example is the influence score function \( \tau^{\text{influence}}_\theta(p) \) which will be considered in Section 4. Another example is the NN score \( \tau^{\text{NN}}_\theta(p) \) that has been studied in our previous work [1], so it will not be examined again in this paper. The condition \( \theta \) is dropped whenever the context is clear.

In this paper, we assume that the object data set \( D \) is indexed by an R-tree and each feature data set \( F_c \) is indexed by an \( \text{MAX} \) aR-tree, where each nonleaf entry augments the maximum quality (of features) in its subtree. Nevertheless, our solutions are directly applicable to data sets that are indexed by other hierarchical spatial indexes (e.g., point quad-trees). The rationale of indexing different feature data sets by separate aR-trees is that: 1) a user queries for only a few features (e.g., restaurants and cafes) out of all possible features (e.g., restaurants, cafes, hospital, market, etc.), and 2) different users may consider different subsets of features.

Based on the above indexing scheme, we develop various algorithms for processing top-k spatial preference queries. Table 1 lists the notations to be used throughout the paper.

### 3.2 Probing Algorithms

We first introduce a brute-force solution that computes the score of every point \( p \in D \) in order to obtain the query results. Then, we propose a group evaluation technique that computes the scores of multiple points concurrently.

#### 3.2.1 Simple Probing Algorithm

According to Section 3.1, the quality \( \omega(s) \) of any feature point \( s \) falls into the interval \([0, 1]\). Thus, for a point \( p \in D \), where not all its component scores are known, its upper bound score \( \tau_\gamma(p) \) is defined as

\[
\tau_\gamma(p) = \sum_{c=1}^{m} \left\{ \begin{array}{ll}
\tau_c(p), & \text{if } \tau_c(p) \text{ is known}, \\
1, & \text{otherwise}.
\end{array} \right.
\] (3)

It is guaranteed that the bound \( \tau_\gamma(p) \) is greater than or equal to the actual score \( \tau(p) \).

Algorithm 1 is a pseudocode of the simple probing (SP) algorithm, which retrieves the query results by computing the score of every object point. The algorithm uses two global variables: \( W_k \) is a min-heap for managing the top-k results and \( \gamma \) represents the top-k score so far (i.e., lowest score in \( W_k \)). Initially, the algorithm is invoked at the root node of the object tree (i.e., \( N = D.root \)). The procedure is recursively applied (at Line 4) on tree nodes until a leaf node is accessed. When a leaf node is reached, the component score \( \tau_c(e) \) (at Line 8) is computed by executing a range search on the feature tree \( F_c \) for range score queries. Lines 6-8 describe an incremental computation technique, for reducing unnecessary component score computations. In particular, the point \( e \) is ignored as soon as its upper bound score \( \tau_c(e) \) (see (3)) cannot be greater than the best-k score \( \gamma \). The variables \( W_k \) and \( \gamma \) are updated when the actual score \( \tau(e) \) is greater than \( \gamma \).

**Algorithm 1. Simple Probing Algorithm**

**algorithm** SP(Node \( N \))

1. **for each** entry \( e \in N \) **do**
2. 2. **if** \( N \) is nonleaf **then**
3. 3. read the child node \( N' \) pointed by \( e \);
4. 4. SP(\( N' \));
5. **else**
6. 6. **for** \( e := 1 \) **to** \( m \) **do**
7. 7. **if** \( \tau_{\gamma}(e) > \gamma \) **then**
8. 8. compute \( \tau_{\gamma}(e) \) using tree \( F_c \); update \( \tau_{\gamma}(e) \);
9. 9. **if** \( \tau(e) > \gamma \) **then**
10. 10. update \( W_k \) (and \( \gamma \)) by \( e \);

#### 3.2.2 Group Probing Algorithm

Due to separate score computations for different objects, SP is inefficient for large-object data sets. In view of this, we propose the group probing (GP) algorithm, a variant of SP, that reduces I/O cost by computing scores of objects in the same leaf node of the R-tree concurrently. In GP, when a leaf node is visited, its points are first stored in a set \( V \) and then their component scores are computed concurrently at a single traversal of the \( F_c \) tree.

We now introduce some distance notations for MBRs. Given a point \( p \) and an MBR \( e \), the value mindist\((p, e)\) (maxdist\((p, e)\)) [4] denotes the minimum (maximum) possible distance between \( p \) and any point in \( e \). Similarly, given two MBRs \( e_a \) and \( e_b \) the value mindist\((e_a, e_b)\)
Like (3), with the component scores on this idea. BB is called with numerous score computations. \(T(e)\) for computing bound score for any point in the subtree of nonleaf entries since the subtree of mindist from some point \(v \in V\) is within the range \(\epsilon\), then the procedure is applied recursively on the child node of \(e\), since the subtree of \(F_c\) rooted at \(e\) may contribute to the component score of \(p\). In case \(e\) is a leaf entry (i.e., a feature point), the scores of points in \(V\) are updated if they are within distance \(\epsilon\) from \(e\).

Algorithm 2. Group Range Score Algorithm

\[\text{algorithm Group\_Range(Node } N, \text{ Set } V, \text{ Value } c, \text{ Value } e)\]

1: for each entry \(e \in N\) do
2: \hspace{1em} if \(N\) is nonleaf then
3: \hspace{2em} if \(\exists p \in V, \text{mindist}(p,e) \leq \epsilon\) then
4: \hspace{3em} read the child node \(N'\) pointed by \(e\);
5: \hspace{3em} Group\_Range\((N',V,e)\);
6: \hspace{2em} else
7: \hspace{3em} for each \(p \in V\) such that \(\text{dist}(p,e) \leq \epsilon\) do
8: \hspace{4em} \(T_e(p) := \max\{T_e(p), \omega(e)\}\);

3.3 Branch-and-Bound Algorithm

GP is still expensive as it examines all objects in \(D\) and computes their component scores. We now propose an algorithm that can significantly reduce the number of objects to be examined. The key idea is to compute, for nonleaf entries \(e\) in the object tree \(D\), an upper bound \(T(e)\) of the score \(T(p)\) for any point \(p\) in the subtree of \(e\). If \(T(e) \leq \gamma\), then we need not access the subtree of \(e\), thus we can save numerous score computations.

Algorithm 3 is a pseudocode of our BB algorithm, based on this idea. BB is called with \(N\) being the root node of \(D\). If \(N\) is a nonleaf node, Lines 3-5 compute the scores \(T(e)\) for nonleaf entries \(e\) concurrently. Recall that \(T(e)\) is an upper bound score for any point in the subtree of \(e\). The techniques for computing \(T(e)\) will be discussed shortly. Like (3), with the component scores \(T_e(e)\) known so far, we can derive \(T_e(c)\), an upper bound of \(T(e)\). If \(T_e(c) \leq \gamma\), then the subtree of \(c\) cannot contain better results than those in \(W_k\) and it is removed from \(V\). In order to obtain points with high scores early, we sort the entries in descending order of \(T(e)\) before invoking the above procedure recursively on the child nodes pointed by the entries in \(V\). If \(N\) is a leaf node, we compute the scores for all points of \(N\) concurrently and then update the set \(W_k\) of the top-\(k\) results. Since both \(W_k\) and \(\gamma\) are global variables, their values are updated during recursive call of BB.

Algorithm 3. Branch-and-Bound Algorithm

\[W_k := \text{new min-heap of size } k \text{ (initially empty)};\]
\[\gamma := 0;\]
\[\triangleright k\text{th score in } W_k\]

\[\text{algorithm BB(Node } N)\]
1: \(V := \{e \mid e \in N\}\);
2: if \(N\) is nonleaf then
3: \hspace{1em} for \(c := 1 \text{ to } m\) do
4: \hspace{2em} compute \(T_e(c)\) for all \(e \in V\) concurrently;
5: \hspace{2em} remove entries \(e \in V\) such that \(T_e(c) \leq \gamma\);
6: \hspace{2em} sort entries \(e \in V\) in descending order of \(T(e)\);
7: \hspace{2em} for each entry \(e \in V\) such that \(T(e) > \gamma\) do
8: \hspace{3em} read the child node \(N'\) pointed by \(e\);
9: \hspace{2em} \(\triangleright e\)th score in \(W_k\);
10: else
11: \hspace{1em} for \(c := 1 \text{ to } m\) do
12: \hspace{2em} compute \(T_e(c)\) for all \(e \in V\) concurrently;
13: \hspace{2em} remove entries \(e \in V\) such that \(T_e(c) \leq \gamma\);
14: \hspace{2em} update \(W_k\) and \(\gamma\) by entries in \(V\);

3.3.1 Upper Bound Score Computation

It remains to clarify how the (upper bound) scores \(T_e(c)\) of nonleaf entries (within the same node \(N\)) can be computed concurrently (at Line 4). Our goal is to compute these upper bound scores such that

- the bounds are computed with low I/O cost, and
- the bounds are reasonably tight, in order to facilitate effective pruning.

To achieve this, we utilize only level-1 entries (i.e., lowest level nonleaf entries) in \(F_c\) for deriving upper bound scores because: 1) there are much fewer level-1 entries than leaf entries (i.e., points), and 2) high-level entries in \(F_c\) cannot provide tight bounds. In our experimental study, we will also verify the effectiveness and the cost of using level-1 entries for upper bound score computation.

Algorithm 2 can be modified for the above upper bound computation task (where input \(V\) corresponds to a set of nonleaf entries), after changing Line 2 to check whether child nodes of \(N\) are above the leaf-level.

The following example illustrates how upper bound range scores are derived. In Fig. 4a, \(v_1\) and \(v_2\) are nonleaf entries in the object tree \(D\) and the others are level-1 entries in the feature tree \(F_c\). For the entry \(v_1\), we first define its Minkowski region [21] (i.e., gray region around \(v_1\)), the area whose mindist from \(v_1\) is within \(\epsilon\). Observe that only entries \(e_i\) intersecting the Minkowski region of \(v_1\) can contribute to the score of some point in \(v_1\). Thus, the upper bound score \(T_e(c)\) is simply the maximum quality of entries \(e_1, e_2, e_3, e_6, e_7\), i.e., 0.9. Similarly, \(T_e(c)\) is computed as the maximum quality of entries \(e_2, e_3, e_4, e_6, e_7\), i.e., 0.7. Assuming that \(v_1\) and \(v_2\) are entries in the same tree node of \(D\), their upper bounds are computed concurrently to reduce I/O cost.
3.4 Optimized Branch-and-Bound Algorithm

This section develops a more efficient score computation technique to reduce the cost of the BB algorithm.

3.4.1 Motivation

Recall that Lines 11-13 of the BB algorithm are used to compute the scores of object points (i.e., leaf entries of the R-tree on D). A leaf entry \( e \) is pruned if its upper bound score \( \tau_e(e) \) is not greater than the best score found so far \( \gamma \). However, the upper bound score \( \tau_e(e) \) (see (3)) is not tight because any unknown component score is replaced by a loose bound (i.e., the value 1).

Let us examine the computation of \( \tau_e(p_1) \) for the point \( p_1 \) in Fig. 4b. The entry \( e_{21}^F \) is a nonleaf entry from the feature tree \( F \). Its augmented quality value is \( \omega(e_{21}^F) = 0.8 \). The entry points to a leaf node containing two feature points, whose qualities values are 0.6 and 0.8, respectively. Similarly, \( e_{22}^F \) is a nonleaf entry from the tree \( F_2 \) and it points to a leaf node of feature points.

Suppose that the best score found so far in BB is \( \gamma = 1.4 \) (not shown in the figure). We need to check whether the score of \( p_1 \) can be higher than \( \gamma \). For this, we compute the first component score \( \tau_1(p_1) = 0.6 \) by accessing the child node of \( e_{21}^F \). Now, we have the upper bound score of \( p_1 \) as \( \tau_{3}(p_1) = 0.6 + 1.0 = 1.6 \). Such a bound is above \( \gamma = 1.4 \) so we need to compute the second component score \( \tau_2(p_1) = 0.5 \) by accessing the child node of \( e_{22}^F \). The exact score of \( p_1 \) is \( \tau(p_1) = 0.6 + 0.5 = 1.1 \); the point \( p_1 \) is then pruned because \( \tau(p_1) \leq \gamma \). In summary, two leaf nodes are accessed during the computation of \( \tau(p_1) \).

Our observation here is that the point \( p_1 \) can be pruned earlier, without accessing the child node of \( e_{22}^F \). By taking the maximum quality of level-1 entries (from \( F_2 \)) that intersect the \( \epsilon \)-range of \( p_1 \), we derive: \( \tau_2(p_1) \leq \omega(e_{22}^F) = 0.7 \). With the first component score \( \tau_1(p_1) = 0.6 \), we infer that: \( \tau(p_1) \leq 0.6 + 0.7 = 1.3 \). Such a value is below \( \gamma \) so \( p_1 \) can be pruned.

3.4.2 Optimized Computation of Scores

Based on our observation, we propose a tighter derivation for the upper bound score of \( p \) than the one shown in (3).

Let \( p \) be an object point in \( D \). Suppose that we have traversed some paths of the feature trees on \( F_1, F_2, \ldots, F_m \). Let \( \mu_e \) be an upper bound of the quality value for any unvisited entry (leaf or nonleaf) of the feature tree \( F_e \). We then define the function \( \tau_e(p) \) as

\[
\tau_e(p) = \max_{c = 1}^m \left\{ \max\left\{ \omega(s) : s \in F_e, \text{dist}(p, s) \leq \epsilon, \omega(s) \geq \mu_c \right\} \right\},
\]

where \( \max \) denotes the maximum upper bound quality of any visited feature point within distance \( \epsilon \) from \( p \).

The following lemma shows that the value \( \tau_e(p) \) is always greater than or equal to the actual score \( \tau(p) \).

**Lemma 1.** It holds that \( \tau_e(p) \geq \tau(p) \), for any \( p \in D \).

**Proof.** If the actual component score \( \tau_e(p) \) is above \( \mu_e \), then, \( \tau_e(p) = \max\{\omega(s) : s \in F_e, \text{dist}(p, s) \leq \epsilon, \omega(s) \geq \mu_e \} \).

Otherwise, we derive \( \tau_e(p) \leq \mu_e \). In both cases, we have \( \tau_e(p) \geq \mu_e \).

Therefore, we have \( \tau_e(p) \geq \tau(p) \).

According to (4), the value \( \tau_e(p) \) is tight only when every \( \mu_e \) value is low. In order to achieve this, we access the feature trees in a round-robin fashion, and traverse the entries in each feature tree in descending order of quality values. Round-robin is a popular and effective strategy used for efficient merging of rankings [7], [9]. Alternative strategies include the selectivity-based strategy and the fractal-dimension strategy [22]. These strategies are designed specifically for coping with high-dimensional data, however, in our problem setting, they have insignificant performance gain over round-robin.

Algorithm 4 is the pseudocode for computing the scores of objects efficiently from the feature trees \( F_1, F_2, \ldots, F_m \). The set \( V \) contains objects whose scores need to be computed. Here, \( \epsilon \) refers to the distance threshold of the range score, and \( \gamma \) represents the best score found so far. For each feature tree \( F_e \), we employ a max-heap \( H_e \) to traverse the entries of \( F_e \) in descending order of their quality values. The root of \( F_e \) is first inserted into \( H_e \). The variable \( \mu_e \) maintains the upper bound quality of entries in the tree that will be visited. We then initialize each component score \( \tau_c(p) \) of every object \( p \in V \) to 0.

**Algorithm 4.** Optimized Group Range Score Algorithm (Trees)

\[
\text{for } e \in C_f, F_2, \ldots, F_m, \text{ Set } V, \text{ Value } \epsilon, \text{ Value } \gamma
\]

1: \( \text{for } e = 1 \text{ to } m \text{ do} \)
2: \( H_e := \text{new max-heap (with quality score as key)} \)
3: \( \text{insert } F_e, \text{root into } H_e \)
4: \( \mu_e := 1 \)
5: \( \text{for each entry } p \in V \text{ do} \)
6: \( \tau_e(p) := 0 \)
7: \( \alpha := 1 \)
8: \( \text{while } |V| > 0 \text{ and there exists a nonempty heap } H_0 \text{ do} \)
9: \( \text{deheap an entry } \epsilon \text{ from } H_0 \)
10: \( \mu_e := \omega(\epsilon) \)
11: \( \text{if } \forall p \in V, \text{mindist}(p, \epsilon) > \epsilon \text{ then} \)
12: \( \text{update threshold} \)
13: \( \text{continue at Line 8} \)
14: \( \text{for each } p \in V \text{ do} \)
15: \( \text{prune unqualified points} \)
16: \( \text{if } (\sum_{c=1}^m \max\{\mu_e, \tau_c(p)\}) \leq \gamma \text{ then} \)
17: \( \text{remove } p \text{ from } V \)
18: \( \text{read the child node } CN \text{ pointed to by } \epsilon \)
19: \( \text{for each entry } e' \text{ of } CN \text{ do} \)
20: \( \text{if } CN \text{ is a nonleaf node then} \)
21: \( \text{if } \exists p \in V, \text{mindist}(p, e') \leq \epsilon \text{ then} \)
22: \( \text{insert } e' \text{ into } H_0 \)
23: \( \text{else} \)
24: \( \text{update component scores} \)
25: \( \text{for each } p \in V \text{ such that } \text{dist}(p, e') \leq \epsilon \text{ do} \)
26: \( \tau_c(p) := \max\{\tau_c(p), \omega(e')\} \)
27: \( \text{for each entry } p \in V \text{ do} \)
28: \( \tau(p) := \sum_{c=1}^m \tau_c(p) \)

At Line 7, the variable \( \alpha \) keeps track of the ID of the current feature tree being processed. The loop at Line 8 is used to compute the scores for the points in the set \( V \). We
then deheap an entry \( e \) from the current heap \( H_o \). The property of the max-heap guarantees that the quality value of any future entry dedequeed from \( H_o \) is at most \( \omega(e) \). Thus, the bound \( \mu_e \) is updated to \( \omega(e) \). At Lines 11-12, we prune the entry \( e \) if its distance from each object point \( p \in V \) is larger than \( \epsilon \). In case \( e \) is not pruned, we compute the tight upper bound score \( \tau_{\rho}(p) \) for each \( p \in V \) (by (4)); the object \( p \) is removed from \( V \) if \( \tau_{\rho}(p) \leq \gamma \) (Lines 13-15).

Next, we access the child node pointed to by \( e \), and examine each entry \( e' \) in the node (Lines 16-17). A nonleaf entry \( e' \) is inserted into the heap \( H_o \) if its minimum distance from some \( p \in V \) is within \( \epsilon \) (Lines 18-20); whereas a leaf entry \( e' \) is used to update the component score \( \tau_{\rho}(p) \) for any \( p \in V \) within distance \( \epsilon \) from \( e' \) (Lines 22-23). At Line 24, we apply the round-robin strategy to find the next \( \alpha \) such that the heap \( H_o \) is not empty. The loop at Line 8 repeats while \( V \) is not empty and there exists a nonempty heap \( H_o \). At the end, the algorithm derives the exact scores for the remaining points of \( V \).

### 3.4.3 The BB* Algorithm

Based on the above, we extend BB (Algorithm 3) to an optimized BB* algorithm as follows: First, Lines 11-13 of BB are replaced by a call to Algorithm 4, for computing the exact scores for object points in the set \( V \). Second, Lines 3-5 of BB are replaced by a call to a modified Algorithm 4, for deriving the upper bound scores for nonleaf entries (in \( V \)). Such a modified Algorithm 4 is obtained after replacing Line 18 by checking whether the node \( CN \) is a nonleaf node above the level-1.

### 3.5 Feature Join Algorithm

An alternative method for evaluating a top-\( k \) spatial preference query is to perform a multway spatial join \cite{23} on the feature trees \( F_1, F_2, \ldots, F_m \) to obtain combinations of feature points which can be in the neighborhood of some object from \( D \). Spatial regions which correspond to combinations of high scores are then examined, in order to find data objects in \( D \) having the corresponding feature combination in their neighborhood. In this section, we first introduce the concept of a combination, then discuss the conditions for a combination to be pruned, and finally elaborate the algorithm used to progressively identify the combinations that correspond to query results.

Tuple \((f_1, f_2, \ldots, f_m)\) is a combination if, for any \( c \in [1, m] \), \( f_c \) is an entry (either leaf or nonleaf) in the feature tree \( F_c \). The score of the combination is defined by

\[
\tau((f_1, f_2, \ldots, f_m)) = \sum_{c=1}^{m} \omega(f_c).
\]

For a nonleaf entry \( f_c \), \( \omega(f_c) \) is the \( \max \) of all feature qualities in its subtree (stored with \( f_c \), since \( F_c \) is an aR-tree). A combination disqualifies the query if

\[
\exists (i \neq j \land i, j \in [1, m]), \text{mindist}(f_i, f_j) > 2\epsilon.
\]

When such a condition holds, it is impossible to have a point in \( D \) whose mindist from \( f_i \) and \( f_j \) are both within \( \epsilon \), respectively. The above validity check acts as a multway join condition that significantly reduces the number of combinations to be examined.

![Fig. 5. Qualified combinations for the join. (a) Nonleaf combination. (b) Leaf combination.](image)

Figs. 5a and 5b illustrate the condition for a nonleaf combination \( \langle A_1, B_2 \rangle \) and a leaf combination \( \langle a_3, b_4 \rangle \), respectively, to be a candidate combination for the query.

Algorithm 5 is a pseudocode of our feature join algorithm.

It employs a max-heap \( H \) for managing combinations of feature entries in descending order of their combination scores. The score of a combination \( \langle f_1, f_2, \ldots, f_m \rangle \) as defined in (5) is an upper bound of the scores of all combinations \( \langle s_1, s_2, \ldots, s_m \rangle \) of feature points, such that \( s_c \) is located in the subtree of \( f_c \) for each \( c \in [1, m] \). Initially, the combination with the root pointers of all feature trees is enheaped. We progressively deheap the combination with the largest score.

If all its entries point to leaf nodes, then we load these nodes \( L_1, \ldots, L_m \) and call \texttt{Find_Result} to traverse the object R-tree \( D \) and find potential results. \texttt{Find_Result} is a variant of the BB algorithm, with the following difference: \( L_1, \ldots, L_m \) are viewed as \( m \) tiny feature trees (each with one node) and accesses to them incur no extra I/O cost.

#### Algorithm 5. Feature Join Algorithm

1. \( W_k := \text{new min-heap of size } k \) (initially empty);
2. \( \gamma := 0; \)
3. for \( k:1 \) to \( m \) do
4. 4. deheap \( \langle f_1, f_2, \ldots, f_m \rangle \) from \( H \);
5. 5. if \( \forall c \in [1, m], f_c \) points to a leaf node
6. 6. 6. for \( c := 1 \) to \( m \) do
7. 7. 7. read the child node \( L_c \) pointed by \( f_c \);
8. 8. 8. \texttt{Find_Result}(\( D \).root, \( L_1, \ldots, L_m \));
9. 9. else
10. 10. 10. \( f_c := \text{highest level entry among } f_1, f_2, \ldots, f_m \);
11. 11. 11. read the child node \( N_c \) pointed by \( f_c \);
12. 12. 12. for each entry \( e_c \in N_c \) do
13. 13. 13. insert \( \langle f_1, f_2, \ldots, e_c, \ldots, f_m \rangle \) into \( H \) if its score is greater than \( \gamma \) and it qualifies the query;
14. end
15. end
16. end
17. end
18. end
19. while \( H \) is not empty do
20. 20. end
21. end
22. \texttt{Find_Result}(\texttt{Node} \( N \), \texttt{Nodes} \( L_1, \ldots, L_m \))
23. 23. end
24. end
25. 25. for each entry \( e \in N \) do
26. 26. if \( N \) is nonleaf then
27. 27. compute \( T(e) \) by entries in \( L_1, \ldots, L_m \);
28. 28. if \( T(e) > \gamma \) then
29. 29. read the child node \( N' \) pointed by \( e \);
30. 30. \texttt{Find_Result}(\( N' \), \( L_1, \ldots, L_m \));
31. 31. else
32. 32. compute \( \tau(e) \) by entries in \( L_1, \ldots, L_m \);
33. 33. update \( W_k \) (and \( \gamma \)) by \( e \) (when necessary);
3.6 Extension to Monotonic Aggregate Functions

We now extend our proposed solutions for processing the top-\(k\) spatial preference query defined by any monotonic aggregate function \(\text{AGG}\). Examples of \(\text{AGG}\) include (but not limited to) the MIN and MAX functions.

3.6.1 Adaptation of Incremental Computation

Recall that the incremental computation technique is applied by algorithms SP, GP, and BB, for reducing I/O cost. Specifically, even if some component score \(\tau_s(p)\) of a point \(p\) has not been computed yet, the upper bound score \(\tau_u(p)\) of \(p\) can be derived by (3). Whenever \(\tau_s(p)\) drops below the best score found so far \(\gamma\), the point \(p\) can be discarded immediately without needing to compute the unknown component scores of \(p\).

In fact, the algorithms SP, GP, and BB are directly applicable to any monotonic aggregate function \(\text{AGG}\) because (3) can be generalized for \(\text{AGG}\). Now, the upper bound score \(\tau_u(p)\) of \(p\) is defined as

\[
\tau_u(p) = \text{AGG} \bigg\{ \tau_s(p) \bigg\}_{s=1}^{m} \quad \text{if} \quad \tau_s(p) \text{ is known, otherwise.}
\]  

(7)

Due to the monotonicity property of \(\text{AGG}\), the bound \(\tau_u(p)\) is guaranteed to be greater than or equal to the actual score \(\tau(p)\).

3.6.2 Adaptation of Upper Bound Computation

The BB* and FJ algorithms compute the upper bound score of a nonleaf entry of the object tree \(D\) or a combination of entries from feature trees, by summing its upper bound component scores. Both BB* and FJ are applicable to any monotonic aggregate function \(\text{AGG}\), with only the slight modifications discussed below. For BB*, we replace the summation operator by \(\text{AGG}\), in (4), and at Lines 14 and 26 of Algorithm 4. For FJ, we replace the summation by \(\text{AGG}\), in (5).

4 Influence Score

This section first studies the influence score function that combines both the qualities and relative locations of feature points. It then presents the adaptations of our solutions in Section 3 for the influence score function. Finally, we discuss how our solutions can be used for other types of influence score functions.

4.1 Score Definition

The range score has a drawback that the parameter \(\epsilon\) is not easy to set. Consider, for instance, the example of the range score \(\tau^\text{rng}\) in Fig. 6a, where the white points are object points in \(D\), the gray points and black points are feature points in the feature sets \(F_1\) and \(F_2\), respectively. If \(\epsilon\) is set to 0.2 (shown by circles), then the object point \(p_2\) has the score \(\tau^\text{rng}(p_2) = 0.9 + 0.1 = 1.0\) and it cannot be the best object (as \(\tau^\text{rng}(p_1) = 1.2\)). This happens because a high-quality black feature is barely outside the \(\epsilon\)-range of \(p_2\). Had \(\epsilon\) been slightly larger, that black feature (with weight 0.6) would contribute to the score of \(p_2\), making it the best object.

In the field of statistics, the Gaussian density function [24] has been used to estimate the density in the space, from a set \(F\) of points. The density at location \(p\) is estimated as:

\[
G(p) = \sum_{f \in F} \exp(-\frac{\text{dist}(p,f)}{2})
\]

where \(\text{dist}(p,f)\) is a parameter. Its advantage is that the value \(G(p)\) is not sensitive to the range \(\epsilon\). \(G(p)\) is mainly contributed by the points (of \(F\)) close to \(p\) and weakly affected by the points far away.

Inspired by the above function, we devise a score function such that it is not too sensitive to the range parameter \(\epsilon\). In addition, the users in our application usually prefer a high-quality restaurant (i.e., a feature point) rather than a large number of low-quality restaurants. Therefore, we use the maximum operator rather than the summation in \(G(p)\). Specifically, we define the influence score of an object point \(p\) with respect to the feature set \(F\), as

\[
\tau^\text{inf}(p) = \max \{ \omega(s) \cdot 2^{-\frac{\text{dist}(p,s)}{\epsilon}} | s \in F \},
\]

where \(\omega(s)\) is the quality of \(s\), \(\epsilon\) is a user-specified range, and \(\text{dist}(p,s)\) is the distance between \(p\) and \(s\).

The overall score \(\tau^\text{inf}(p)\) of \(p\) is then defined as

\[
\tau^\text{inf}(p) = \text{AGG} \{ \tau^\text{inf}(p) | c \in [1, m] \},
\]

where \(\text{AGG}\) is a monotone aggregate operator and \(m\) is the number of feature data sets. Again, we focus on the case where \(\text{AGG}\) is the SUM function.

Let us compute the influence score \(\tau^\text{inf}\) for the points in Fig. 6a, assuming \(\epsilon = 0.2\). From Fig. 6a, we obtain

\[
\tau^\text{inf}(p_1) = \max \{ 0.7 \cdot 2^{-\frac{0.2}{0.2}}, 0.9 \cdot 2^{-\frac{0.2}{0.2}} \} = 0.643
\]

and

\[
\tau^\text{inf}(p_2) = \max \{ 0.9 \cdot 2^{-\frac{0.2}{0.2}}, 0.7 \cdot 2^{-\frac{0.2}{0.2}} \} = 0.762.
\]

The top-1 point is \(p_2\), implying that the influence score can capture feature points outside the range \(\epsilon = 0.2\). In fact, the influence score function possesses two nice properties. First, a feature point \(s\) that is barely outside the range \(\epsilon\) (from the object point \(p\)) still has potential to contribute to the score, provided that its quality \(\omega(s)\) is sufficiently high. Second, the distance \(\text{dist}(p,s)\) has an exponentially decaying effect.
on the score, meaning that feature points nearer to \( p \) contribute higher scores.

4.2 Query Processing for SP, GP, BB, and BB*

We now examine the extensions of the SP, GP, BB, and BB* algorithms for top-\( k \) spatial preference queries defined by the influence score in (8).

4.2.1 Incremental Computation Technique

Observe that the upper bound of \( \tau_{e}(p) \) is 1. Therefore, (3) still holds for the influence score, and the incremental computation technique (see Section 3.2) can still be applied in SP, GP, and BB.

4.2.2 Exact Score Computation for a Single Object

For the SP algorithm, we elaborate how to compute the score \( \tau_{e}(p) \) (see (8)) of an object \( p \in \mathcal{D} \). This is challenging because some feature \( s \in \mathcal{F}_{e} \) outside the \( \epsilon \)-range of \( p \) may contribute to the score. Unlike the computation of the range score, we can no longer use the \( \epsilon \)-range to restrict the search space.

Given an object point \( p \) and an entry \( e \) from the feature tree of \( \mathcal{F}_{e} \), we define the upper bound function

\[
\omega^{\text{inf}}(e, p) = \omega(e) \cdot 2^{-\mindist(e, p)}.
\]

In case \( e \) is a leaf entry (i.e., a feature point \( s \)), we have \( \omega^{\text{inf}}(s, p) = \omega(s) \cdot 2^{-\mindist(e, s)} \). The following lemma shows that the value \( \omega^{\text{inf}}(e, p) \) is an upper bound of \( \omega^{\text{inf}}(e', p) \) for any entry \( e' \) in the subtree of \( e \).

**Lemma 2.** Let \( e \) and \( e' \) be entries from the feature tree \( \mathcal{F}_{e} \), such that \( e' \) is in the subtree of \( e \). It holds that \( \omega^{\text{inf}}(e, p) \geq \omega^{\text{inf}}(e', p) \), for any object point \( p \in \mathcal{D} \).

**Proof.** Let \( p \) be any object point \( p \in \mathcal{D} \). Since \( e' \) falls into the subtree of \( e \), we have: \( \mindist(p, e) \leq \mindist(p, e') \). As \( \mathcal{F}_{e} \) is a \( \text{MAX} \) \( \text{R} \)-tree, we have: \( \omega(e) \leq \omega(e') \). Thus, we have: \( \omega^{\text{inf}}(e, p) \geq \omega^{\text{inf}}(e', p) \). \(\square\)

As an example, Fig. 6b shows an object point \( p \) and three entries \( e_1, e_2, e_3 \) from the same feature tree. Note that \( e_2 \) and \( e_3 \) are in the subtree of \( e_1 \). The dotted lines indicate the minimum distance from \( p \) to \( e_1 \) and \( e_3 \), respectively. Thus, we have \( \omega^{\text{inf}}(e_1, p) = 0.8 \cdot 2^{-0.5} = 0.4 \) and \( \omega^{\text{inf}}(e_3, p) = 0.7 \cdot 2^{-0.5} = 0.247 \). Clearly, \( \omega^{\text{inf}}(e_1, p) \) is larger than \( \omega^{\text{inf}}(e_3, p) \).

By using Lemma 2, we apply the best-first approach to compute the exact component score \( \tau_{e}(p) \) for the point \( p \); Algorithm 6 employs a max-heap \( H \) in order to visit the entries of the tree in descending order of their \( \omega^{\text{inf}} \) values. We first insert the root of the tree \( \mathcal{F}_{e} \) into \( H \), and initialize \( \tau_{e}(p) \) to 0. The loop at Line 4 continues as long as \( H \) is not empty. At Line 5, we dequeue an entry \( e \) from \( H \). If the value \( \omega^{\text{inf}}(e, p) \) is above the current \( \tau_{e}(p) \), then there is potential to update \( \tau_{e}(p) \) by using some point in the subtree of \( e \). In that case, we read the child node pointed to by \( e \) and examine each entry \( e' \) in that node (Lines 7-8). If \( e' \) is a nonleaf entry, it is inserted into \( H \) provided that its \( \omega^{\text{inf}}(e', p) \) value is above \( \tau_{e}(p) \). Otherwise, it is used to update \( \tau_{e}(p) \).

**Algorithm 6.** Object Influence Score Algorithm

```plaintext
1: \( H := \text{new max-heap with } \omega^{\text{inf}} \text{ value as key} \)
2: insert \( (\mathcal{F}_{e}, \text{root}, 1,0) \) into \( H \)
3: \( \tau_{e}(p) := 0 \)
4: while \( H \) is not empty do
5: deheap an entry \( e \) from \( H \)
6: if \( \omega^{\text{inf}}(e, p) > \tau_{e}(p) \) then
7: read the child node \( CN \) pointed to by \( e \)
8: for each entry \( e' \) of \( CN \) do
9: if \( CN \) is a nonleaf node then
10: insert \( (e', \omega^{\text{inf}}(e', p)) \) into \( H \)
11: \( \tau_{e}(p) := \max\{\tau_{e}(p), \omega^{\text{inf}}(e', p)\} \)
12: else \( \tau_{e}(p) := \omega^{\text{inf}}(e', p) \)
13: end if
14: end for
15: end if
16: end while
```

4.2.3 Group Computation and Upper Bound Computation

Recall that, for the case of range scores, both the GP and BB algorithms apply the group computation technique (Algorithm 2) for concurrently computing the component score \( \tau_{e}(p) \) for every object point \( p \) in a given set \( V \). Now, Algorithm 6 can be modified as follows to support concurrent computation of influence scores. First, the parameter \( p \) is replaced by a set \( V \) of objects. Second, we initialize the value \( \tau_{e}(p) \) for each object \( p \in V \) at Line 3 and perform the score update for each \( p \) in \( V \) at Line 13. Third, the conditions at Lines 6 and 10 are checked whether they are satisfied by some object \( p \in V \).

In addition, the BB algorithm (see Algorithm 3) needs to compute the upper bound component score \( T_{e}(c) \) for all nonleaf entries in the current node simultaneously. Again, Algorithm 6 can be modified for this purpose.

4.2.4 Optimized Computation of Scores in BB*

Given an entry \( e \) (from a feature tree), we define the upper bound score of \( e \) using a set \( V \) of points as

\[
\omega^{\text{inf}}(e, V) = \max_{p \in V} \omega^{\text{inf}}(e, p).
\]

The BB* algorithm applies Algorithm 4 to compute the range scores for a set \( V \) of object points. With (11), we can modify Algorithm 4 to compute the influence score, with the following changes. First, the heap \( H \) (at Line 2) is used to organize its entries \( e \) in descending order of the key \( \omega^{\text{inf}}(e, V) \), and the value \( \omega(e) \) (at Line 10) is replaced by \( \omega^{\text{inf}}(e, V) \). Second, the restrictions based on the \( \epsilon \)-range (at Lines 11-12, 19, and 22) are removed. Third, the value \( \omega(e) \) (at Line 23) needs to be replaced by \( \omega^{\text{inf}}(e', p) \).

4.3 Query Processing for FJ

The FJ algorithm can be adapted for the influence score, but with two changes. Recall that the tuple \( \langle f_1, f_2, \ldots, f_m \rangle \) is said to be a combination if \( f_1 \) is an entry in the feature tree \( \mathcal{F}_{e} \) for any \( c \in [1, m] \).

First, (6) can no longer be used to prune a combination based on distances among the entries in the combination. Any possible combination must be considered if its upper bound score is above the best score found so far \( \gamma \).

Second, (5) is now a loose upper bound value for the influence score because it ignores the distances among the
entries \( f_c \). Therefore, we need to develop a tighter upper bound for the influence score.

The following lemma shows that, given a set \( \Phi \) of rectangles that partition the spatial domain \( \text{DOM} \), the value \( \max_{r \in \Phi} \sum_{c=1}^{m} \omega^\Phi(f_c, r) \) is an upper bound of the value \( \sum_{c=1}^{m} \omega^\Phi(f_c, p) \) for any point \( p \) (in \( \text{DOM} \)).

**Lemma 3.** Let \( \Phi \) be a set of rectangles which partition the spatial domain \( \text{DOM} \). Given the tree entries \( f_1, f_2, \ldots, f_m \) (from the respective feature trees), it holds that \( \max_{r \in \Phi} \sum_{c=1}^{m} \omega^\Phi(f_c, r) \geq \sum_{c=1}^{m} \omega^\Phi(f_c, p) \), for any point \( p \in \text{DOM} \).

**Proof.** Let \( p \) be a point in \( \text{DOM} \). There exists an rectangle \( r' \in \Phi \) such that \( p \) falls into \( r' \). Thus, for any \( c \in [1, m] \), we have \( \text{mindist}(f_c, r') \leq \text{mindist}(f_c, p) \), and derive \( \omega^\Phi(f_c, r') \geq \omega^\Phi(f_c, p) \). By summing all components, we obtain \( \sum_{c=1}^{m} \omega^\Phi(f_c, r') \geq \sum_{c=1}^{m} \omega^\Phi(f_c, p) \). As \( r' \in \Phi \), we also have \( \max_{r \in \Phi} \sum_{c=1}^{m} \omega^\Phi(f_c, r) \geq \sum_{c=1}^{m} \omega^\Phi(f_c, r') \). Therefore, the lemma is proved.

In fact, the above upper bound value

\[
\max_{r \in \Phi} \sum_{c=1}^{m} \omega^\Phi(f_c, r)
\]

can be tightened by dividing the rectangles of \( \Phi \) into smaller rectangles.

Fig. 7a shows the combination \( \langle f_1, f_2 \rangle \), whose entries belong to the feature trees \( F_1 \) and \( F_2 \), respectively. We first partition the domain space into four rectangles \( r_1, r_2, r_3, r_4 \), and then compute their upper bound values (shown in the figure). Thus, the current upper bound score of \( \langle f_1, f_2 \rangle \) is taken as: \( \max\{1.5, 1.0, 0.1, 0.8\} = 1.5 \). To tighten the upper bound, we pick the rectangle \( r_1 \) with the highest value and partition it into four rectangles \( r_{11}, r_{12}, r_{13}, r_{14} \) (see Fig. 7b). Now, the upper bound score of \( \langle f_1, f_2 \rangle \) becomes: \( \max\{0.8, 1.1, 0.8, 0.7, 1.0, 0.1, 0.8\} = 1.1 \). By applying this method iteratively, the upper bound score can be gradually tightened.

Algorithm 7 is the pseudocode for computing the upper bound score for the combination \( \langle f_1, f_2, \ldots, f_m \rangle \) of feature entries. The parameter \( \gamma \) represents the best score found so far (in FJ). The value \( I_{\text{max}} \) is used to control the number of iterations in the algorithm; its typical value is 20. At Line 1, we employ a max-heap \( H \) to organize its rectangles in descending order of their upper bound scores. Then, we insert the spatial domain rectangle into \( H \). The loop at Line 4 continues while \( H \) is not empty and \( I_{\text{max}} > 0 \). After deheaping a rectangle \( r \) from \( H \) (Line 5), we partition it into four child rectangles. Each child rectangle \( r' \) is inserted into \( H \) if its upper bound score is above \( \gamma \). We then decrement \( I_{\text{max}} \) (at Line 10). At the end (Lines 11-14), if the heap \( H \) is not empty, then algorithm returns the key value of \( H \)'s top entry as the upper bound score. Such a value is guaranteed to be the maximum upper bound value in the heap. Otherwise (i.e., empty \( H \)), the algorithm returns \( \gamma \) as the upper bound score because all rectangles with score below \( \gamma \) have been pruned.

**Algorithm 7.** FJ Upper Bound Computation Algorithm

\[
\text{algorithm FJ}\_\text{Influence}(Value \epsilon, Value \gamma, Value I_{\text{max}})
\]

1: \( H := \text{new heap with upper bound score as key} \);
2: let \( \text{DOM} \) be the rectangle of the spatial domain;
3: insert \( \langle \text{DOM}, \sum_{c=1}^{m} \omega(f_c) \rangle \) into \( H \);
4: while \( I_{\text{max}} > 0 \) and \( H \) is not empty do
5: doheap a rectangle \( r \) from \( H \);
6: partition \( r \) into four child rectangles;
7: for each child rectangle \( r' \) do
8: if \( \sum_{c=1}^{m} \omega^\Phi(f_c, r') > \gamma \) then
9: insert \( \langle r', \sum_{c=1}^{m} \omega^\Phi(f_c, r') \rangle \) into \( H \);
10: \( I_{\text{max}} := I_{\text{max}} - 1 \);
11: if \( H \) is not empty then
12: return the key value of the top entry of \( H \);
13: else
14: return \( \gamma \);

### 4.4 Extension to Generic Influence Scores

Our algorithms can be also applied to other types of influence score functions. Given a function \( U: \mathbb{R} \rightarrow \mathbb{R} \), we model a score function as \( \omega^\Phi(e, p) = \omega(e) \cdot U(\text{dist}(p, s)) \), where \( p \) is an object point and \( s \in F_e \) is a feature point.

Let \( e \) be a feature tree entry. The crux of our solution is to redefine the upper bound function \( \omega^\Phi(e, p) \) (like in (10)) such that \( \omega^\Phi(e, p) \geq \omega^\Phi(s, p) \), for any feature point \( s \) in the subtree of \( e \).

In fact, the upper bound function can be expressed as \( \omega^\Phi(e, p) = \omega(e) \cdot U(d) \), where \( d \) is a distance value. Observe that \( d \) must fall in the interval \([\text{mindist}(p, e), \text{maxdist}(p, e)]\). Thus, we apply a numerical method (e.g., the bisection method) to find the value \( d \in [\text{mindist}(p, e), \text{maxdist}(p, e)] \) that maximizes the value of \( U \).

For the special case that \( U \) is a monotonic decreasing function, we can simply set \( d = \text{mindist}(p, e) \) because it definitely maximizes the value of \( U \).

### 5 Experimental Evaluation

In this section, we compare the efficiency of the proposed algorithms using real and synthetic data sets. Each data set is indexed by an aR-tree with 4 K bytes page size. We used an LRU memory buffer whose default size is set to 0.5 percent of the sum of tree sizes (for the object and feature trees used).

Our algorithms were implemented in C++ and experiments were run on a Pentium D 2.8 GHz PC with 1 GB of RAM. In all experiments, we measure both the I/O cost (in number of page faults) and the total execution time (in seconds) of our algorithms. Section 5.1 describes the experimental settings. Sections 5.2 and 5.3 study the performance of the proposed algorithms for queries with range scores and influence scores, respectively. We then present our experimental findings on real data in Section 5.4.
TABLE 2
Range of Parameter Values

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aggregate function</td>
<td>SUM, MIN, MAX</td>
</tr>
<tr>
<td>Buffer size (%)</td>
<td>0.1, 0.2, 0.5, 1, 2, 5, 10</td>
</tr>
<tr>
<td>Object data size, (</td>
<td>D</td>
</tr>
<tr>
<td>Feature data size, (</td>
<td>F</td>
</tr>
<tr>
<td>Number of results, (k)</td>
<td>1, 2, 4, 8, 16, 32, 64</td>
</tr>
<tr>
<td>Number of features, (m)</td>
<td>1, 2, 3, 4, 5</td>
</tr>
<tr>
<td>Query range, (\epsilon)</td>
<td>10, 20, 50, 100, 200</td>
</tr>
</tbody>
</table>

5.1 Experimental Settings
We used both real and synthetic data for the experiments. The real data sets will be described in Section 5.4. For each synthetic data set, the coordinates of points are random values uniformly and independently generated for different dimensions. By default, an object data set contains 200K points and a feature data set contains 100K points. The point coordinates of all data sets are normalized to the 2D space \([0, 10000]^2\).

For a feature data set \(F_c\), we generated qualities for its points such that they simulate a real-world scenario: facilities close to (far from) a town center often have high (low) quality. For this, a single anchor point \(s_0\) is selected such that its neighborhood region contains high number of points. Let \(\text{dist}_{\text{min}}\) (\(\text{dist}_{\text{max}}\)) be the minimum (maximum) distance of a point in \(F_c\) from the anchor \(s_0\). Then, the quality of a feature point \(s\) is generated as

\[
\omega(s) = \left(\frac{\text{dist}_{\text{max}} - \text{dist}(s, s_0)}{\text{dist}_{\text{max}} - \text{dist}_{\text{min}}}\right)^\vartheta,
\]

where \(\text{dist}(s, s_0)\) stands for the distance between \(s\) and \(s_0\), and \(\vartheta\) controls the skewness (default: 1.0) of quality distribution. In this way, the qualities of points in \(F_c\) lie in \([0, 1]\) and the points closer to the anchor have higher qualities. Also, the quality distribution is highly skewed for large values of \(\vartheta\).

We study the performance of our algorithms with respect to various parameters, which are displayed in Table 2 (their default values are shown in bold). In each experiment, only one parameter varies while the others are fixed to their default values.

5.2 Performance on Queries with Range Scores
This section studies the performance of our algorithms for top-\(k\) spatial preference queries on range scores.

Table 3 shows the I/O cost and execution time of the algorithms, for different aggregate functions (SUM, MIN, and MAX). GP has lower cost than SP because GP computes the scores of points within the same leaf node concurrently. The incremental computation technique (used by SP and GP) derives a tight upper bound score (of each point) for the MIN function, a partially tight bound for SUM, and a loose bound for MAX (see Section 3.6). This explains the performance of SP and GP across different aggregate functions. However, the cost of the other methods are mainly influenced by the effectiveness of pruning. BB employs an effective technique to prune unqualified nonleaf entries in the object tree so it outperforms GP. The optimized score computation method enables BB* to save on average 20 percent I/O and 30 percent time of BB. FJ outperforms its competitors as it discovers qualified combination of feature entries early.

We ignore SP in subsequent experiments, and compare the cost of the remaining algorithms on synthetic data sets with respect to different parameters.

Next, we empirically justify the choice of using level-1 entries of feature trees \(F_c\) for the upper bound score computation routine in the BB algorithm (see Section 3.3). In this experiment, we use the default parameter setting and study how the number of node accesses of BB is affected by the level of \(F_c\) used. Table 4 shows the decomposition of node accesses over the tree \(D\) and the trees \(F_c\), and the statistics of upper bound score computation. Each accessed nonleaf node of \(D\) invokes a call of the upper bound score computation routine.

When level-0 entries of \(F_c\) are used, each upper bound computation call incurs a high number (617.5) of node accesses (of \(F_c\)). On the other hand, using level-2 entries for upper bound computation leads to very loose bounds, making it difficult to prune the leaf nodes of \(D\). Observe that the total cost is minimized when level-1 entries (of \(F_c\)) are used. In that case, the node accesses per upper bound computation call is low (15), and yet the obtained bounds are tight enough for pruning most leaf nodes of \(D\).

Fig. 8 plots the cost of the algorithms as a function of the buffer size. As the buffer size increases, the I/O of all algorithms drops. FJ remains the best method, BB* the second, and BB the third; all of them outperform GP by a wide margin. Since the buffer size does not affect the pruning effectiveness of the algorithms, it has a small impact on the execution time.

Fig. 9 compares the cost of the algorithms with respect to the object data size \(|D|\). Since the cost of FJ is dominated by the cost of joining feature data sets, it is insensitive to \(|D|\). On the other hand, the cost of the other methods (GP, BB, and BB*) increases with \(|D|\), as score computations need to be done for more objects in \(D\).

TABLE 3
Effect of the Aggregate Function, Range Scores

<table>
<thead>
<tr>
<th>SUM function</th>
<th>SP</th>
<th>GP</th>
<th>BB</th>
<th>BB*</th>
<th>FJ</th>
</tr>
</thead>
<tbody>
<tr>
<td>I/O</td>
<td>359027</td>
<td>22594</td>
<td>2033</td>
<td>1535</td>
<td>489</td>
</tr>
<tr>
<td>Time (s)</td>
<td>635.0</td>
<td>32.7</td>
<td>3.0</td>
<td>2.0</td>
<td>1.3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>MIN function</th>
<th>SP</th>
<th>GP</th>
<th>BB</th>
<th>BB*</th>
<th>FJ</th>
</tr>
</thead>
<tbody>
<tr>
<td>I/O</td>
<td>215602</td>
<td>16254</td>
<td>611</td>
<td>615</td>
<td>47</td>
</tr>
<tr>
<td>Time (s)</td>
<td>426.8</td>
<td>22.7</td>
<td>0.9</td>
<td>0.8</td>
<td>0.2</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>MAX function</th>
<th>SP</th>
<th>GP</th>
<th>BB</th>
<th>BB*</th>
<th>FJ</th>
</tr>
</thead>
<tbody>
<tr>
<td>I/O</td>
<td>402704</td>
<td>26126</td>
<td>228</td>
<td>106</td>
<td>8</td>
</tr>
<tr>
<td>Time (s)</td>
<td>742.8</td>
<td>38.2</td>
<td>0.3</td>
<td>0.2</td>
<td>0.1</td>
</tr>
</tbody>
</table>

TABLE 4
Effect of the Level of \(F_c\), Used for Upper Bound Score Computation in the BB Algorithm

<table>
<thead>
<tr>
<th>Level</th>
<th>Node accesses (NA)</th>
<th>Upper bound score computation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>of (D) of (F_c)</td>
<td># of calls</td>
</tr>
<tr>
<td>0</td>
<td>3350</td>
<td>53</td>
</tr>
<tr>
<td>1</td>
<td>2365</td>
<td>130</td>
</tr>
<tr>
<td>2</td>
<td>13666</td>
<td>930</td>
</tr>
</tbody>
</table>
Fig. 10 plots the I/O cost of the algorithms with respect to the feature data size $|F|$ (of each feature data set). As $|F|$ increases, the cost of GP, BB, and FJ increases. In contrast, BB* experiences a slight cost reduction as its optimized score computation method (for objects and nonleaf entries) is able to perform pruning early at a large $|F|$ value.

Fig. 11 plots the cost of the algorithms with respect to the number $m$ of feature data sets. The costs of GP, BB, and BB* rise linearly as $m$ increases because the number of component score computations is at most linear to $m$. On the other hand, the cost of FJ increases significantly with $m$, because the number of qualified combinations of entries is exponential to $m$.

Fig. 12 shows the cost of the algorithms as a function of the number $k$ of requested results. GP, BB, and BB* compute the scores of objects in $D$ in batches, so their performance is insensitive to $k$. As $k$ increases, FJ has weaker pruning power and its cost increases slightly.

Fig. 13 shows the cost of the algorithms, when varying the query range $\epsilon$. As $\epsilon$ increases, all methods access more nodes in feature trees to compute the scores of the points. The difference in execution time between BB* and FJ shrinks as $\epsilon$ increases. In summary, although FJ is the clear winner in most of the experimental instances, its performance is significantly affected by the number $m$ of feature data sets. BB* is the most robust algorithm to parameter changes and it is recommended for problems with large $m$.

5.3 Performance on Queries with Influence Scores

We proceed to examine the cost of our algorithms for top-$k$ spatial preference queries on influence scores.

Fig. 14 compares the cost of the algorithms with respect to the number $m$ of feature data sets. The cost follows the trend in Fig. 11. Again, the number of combinations examined by FJ increases exponentially with $m$ so its cost increases rapidly.

Fig. 15 plots the cost of the algorithms by varying the number $k$ of requested results. Observe that FJ becomes more expensive than BB* (in both I/O and time) when the value of $k$ is beyond 8. This is attributed to two reasons. First, FJ incurs extra computational cost as it needs to invoke Algorithm 7 for computing the upper bound score of a combination of feature entries. Second, FJ incurs high I/O cost to identify objects in $D$ that produce high scores with the current combination of features.

Fig. 16 shows the cost of the algorithms as a function of the parameter $\epsilon$. Interestingly, the trend here is different from the one in Fig. 13. According to (8), when $\epsilon$ decreases, the influence score also decreases, rendering it more difficult to distinguish the scores among different objects. Thus, the cost of BB, BB*, and FJ becomes high at a low $\epsilon$ value. Summing up, for the newly introduced influence score, FJ is more sensitive to parameter changes and it loses to BB* not only when there are multiple feature data sets, but also at large $k$. 
5.4 Results on Real Data

In this section, we conduct experiments on real object and feature data sets in order to demonstrate the application of top-$k$ spatial preference queries.

We obtained three real spatial data sets from a travel portal website, http://www.allstays.com/. Locations in these data sets correspond to (longitude and latitude) coordinates in US. We cleaned the data sets by discarding records without longitude and latitude. Each remaining location is normalized to a point in the 2D space $\left[0, 10,000\right]/C138^2$. One data set is used as the object data set $D$ contains 11,399 camping locations. The feature data set $F_1$ contains 30,921 hotel records, each with a room price (quality) and a location. The feature data set $F_2$ has 3,848 records of Wal-Mart stores, each with a gasoline availability (quality) and a location. The domain of each quality attribute (e.g., room price and gasoline availability) is normalized to the unit interval $[0, 1]$. Intuitively, a camping location is considered as good if it is close to a Wal-Mart store with high gasoline availability (i.e., convenient supply) and a hotel with high room price (which indirectly reflects the quality of nearby outdoor environment).

Fig. 17 plots the cost of the algorithms with respect to $\epsilon$, for queries with range scores. At a very small $\epsilon$ value, most of the objects have the zero score as they have no feature points within their neighborhood. This forces BB, BB*, and FJ to access a larger number of objects (or feature combinations) before finding an object with nonzero score, which can then be used for pruning other unqualified objects.

Fig. 18 compares the cost of the algorithms with respect to $\epsilon$, for queries with influence scores. In general, the cost follows the trend in Fig. 16. BB* outperforms BB at low $\epsilon$ value whereas BB incurs a slightly lower cost than BB* at a high $\epsilon$ value. Observe that the cost of BB and BB* is close to that of FJ when $\epsilon$ is sufficiently high. In summary, the relative performance between the algorithms in all experiments is consistent to the results on synthetic data.

6 Conclusion

In this paper, we studied top-$k$ spatial preference queries, which provide a novel type of ranking for spatial objects based on qualities of features in their neighborhood. The neighborhood of an object $p$ is captured by the scoring function: 1) the range score restricts the neighborhood to a crisp region centered at $p$, whereas 2) the influence score relaxes the neighborhood to the whole space and assigns higher weights to locations closer to $p$.

We presented five algorithms for processing top-$k$ spatial preference queries. The baseline algorithm SP computes the scores of every object by querying on feature data sets. The algorithm GP is a variant of SP that reduces I/O cost by computing scores of objects in the same leaf node concurrently. The algorithm BB derives upper bound scores for nonleaf entries in the object tree, and prunes those that cannot lead to better results. The algorithm BB* is a variant of BB that utilizes an optimized method for computing the scores of objects (and upper bound scores of nonleaf entries). The algorithm FJ performs a multiway join on feature trees to obtain qualified combinations of feature points and then search for their relevant objects in the object tree.

Based on our experimental findings, BB* is scalable to large data sets and it is the most robust algorithm with respect to various parameters. However, FJ is the best algorithm in cases where the number $m$ of feature data sets is low and each feature data set is small.
In the future, we will study the top-k spatial preference query on a road network, in which the distance between two points is defined by their shortest path distance rather than their euclidean distance. The challenge is to develop alternative methods for computing the upper bound scores for a group of points on a road network.

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REFERENCES


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