

Reverse Nearest Neighbors in Large Graphs

Man Lung Yiu

Department of Computer Science and Information Systems
University of Hong Kong
Pokfulam Road, Hong Kong
mlyiu2@csis.hku.hk

Nikos Mamoulis

Department of Computer Science and Information Systems
University of Hong Kong
Pokfulam Road, Hong Kong
nikos@csis.hku.hk

Dimitris Papadias

Department of Computer Science
Hong Kong University of Science and Technology
Clearwater Bay, Hong Kong
dimitris@cs.ust.hk

Yufei Tao

Department of Computer Science
City University of Hong Kong
Tat Chee Avenue, Hong Kong
taoyf@cityu.edu.hk

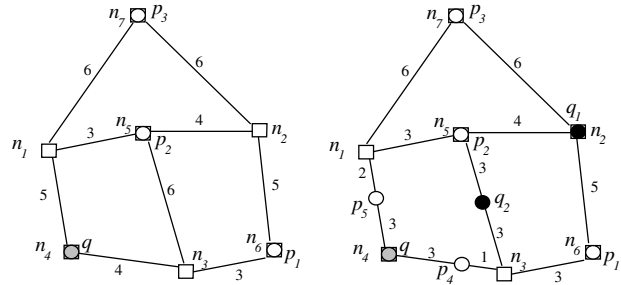
Abstract

A reverse nearest neighbor query returns the data objects that have a query point as their nearest neighbor. Although such queries have been studied quite extensively in Euclidean spaces, there is no previous work in the context of large graphs. In this paper, we propose algorithms and optimization techniques for RNN queries by utilizing some characteristics of networks.

Introduction-Motivation

Given a multi-dimensional dataset P and a point q , a (monochromatic) reverse nearest neighbor (RNN) query retrieves all the points $p \in P$ that have q as their nearest neighbor. Given two datasets P and Q and a point q , a bichromatic (bRNN) query retrieves all the points $p \in P$ that are closer to q than to any point of Q . The problem has received considerable attention the last few years [KM00, SRAA01, TPL04] due to its relevance in several applications involving decision support, resource allocation, profile-based marketing, etc. However, all the existing work focuses exclusively on Euclidean spaces, whereas in several domains the data are represented as large disk-based graphs.

As an example, consider a P2P network, where each point (i.e., peer) $p \in P$ lies on a node $n \in V$, but some nodes may not contain relevant peers to a specific query. We call such graphs *restricted*. For instance, in Figure 1a, assume that a new user q interested in music enters the system. A (monochromatic) RNN query retrieves among the existing users also interested in music (p_1 to p_3), the ones for which q will become their new NN. On the other hand, nodes such as n_1 and n_2 are irrelevant to the user (e.g., they may represent peers with other types of content) and they are considered empty. Given the edge costs of Figure 1a, $\text{RNN}(q) = \{p_1, p_2\}$, i.e., q is most beneficial to p_1 and p_2 since it is their closest NN in terms of network cost and shares the same interests.



(a) RNN in restricted graph (b) bRNN in unrestricted graph
Figure 1 Examples of RNN queries in graphs

In *unrestricted* networks, the query and the data points can reside anywhere on the edges of the graph. Figure 1b shows an example of a bichromatic query in a road network, where points p_1 to p_5 stand for residential blocks and q_1, q_2 indicate restaurants. Nodes n_1 and n_3 are empty road junctions (i.e., they do not contain residential blocks or restaurants). Given several choices for the location of a new restaurant, the query $\text{bRNN}(q)$ may be used to evaluate the benefit of q in terms of the customers that it may attract from rival restaurants based on proximity. Specifically, the set $\text{bRNN}(q) = \{p_4, p_5\}$ represents the blocks that are closer to q than to any other competitor. Similarly, $\text{bRNN}(q_1) = \{p_1, p_3\}$ and $\text{bRNN}(q_2) = \{p_2\}$.

Algorithms-Optimizations

We propose algorithms for monochromatic and bichromatic queries involving an arbitrary number k of RNNs in restricted or unrestricted graphs. Due to space constraints, in the following description we assume restricted networks and that $k=1$. Our algorithms use the following lemma for pruning the search space.

Lemma 1 Let q be a query point, n a graph node and p a data point satisfying $d(q,n) > d(p,n)$. For any point $p' \neq p$ whose shortest path to q passes through n , it holds that $d(q,p') > d(p,p')$, i.e., $p' \notin \text{RNN}(q)$.

Proof. $d(q,p) = d(q,n) + d(n,p) > d(p,n) + d(n,p) \geq d(p,p)$. \square

For instance, in Figure 1a, $d(q,n_3)=4 > d(p_1,n_3)=3$. Thus, any data point (other than p_1) whose shortest path to q passes from n_3 cannot be a RNN of q because it is closer to p_1 (than q). On the other hand, $p_1 \in \text{RNN}(q)$ if there is not other data point within distance $d(p_1,q)$ from p_1 (which is true in this example).

The first algorithm, called *eager*, initializes a heap H and inserts the source node. When a node n is de-heaped, *eager* applies Lemma 1 in order to determine whether the expansion should proceed. In particular, it first retrieves the NN of n by expanding the network around n (we use a technique similar to Dijkstra's algorithm). If no data point is discovered within distance $d(n,q)$ from n , the algorithm en-heaps the adjacent nodes of n . If there is a point p , such that $d(n,q) > d(n,p)$, the expansion does not proceed further because (according to Lemma 1) n cannot lead to a RNN of q . In this case, however, we need to verify (by another network expansion around p) if $p \in \text{RNN}(q)$, in which case p is added to the result. Furthermore, p is marked as *verified* in order not to be expanded, if it is found again in the future through another node.

As an example consider the RNN query of Figure 1a initiated at node n_4 , which is inserted into $H = \langle n_4, 0 \rangle$. Then, n_4 is de-heaped and its adjacent nodes are added to $H = \langle n_3, 4 \rangle, \langle n_1, 5 \rangle$. The subsequent removal (from H) of n_3 triggers an expansion around n_3 for finding potential data points closer than the query. In this case, $d(n_3, p_1) = 3 < d(n_3, q)$, meaning (by Lemma 1) that we do not search farther, i.e., the adjacent nodes of n_3 are not inserted to H . Since $q = \text{NN}(p_1)$, p_1 is an actual result. Next, n_1 is de-heaped, its NN p_2 is discovered and the search for RNNs terminates because $d(n_1, q) = 5 > d(n_1, p_2) = 3$. The final step simply verifies p_2 as an actual result because $q = \text{NN}(p_2)$.

Although *eager* minimizes the number of nodes inserted into the heap, it may perform numerous local network expansions for (i) retrieving the nearest point p of a de-heaped node n and (ii) for verifying whether $p \in \text{RNN}(q)$. The *lazy* algorithm delays pruning until a point is visited. In particular, assuming single RNN retrieval, when the de-heaped node n does not contain a point, *lazy* simply inserts its adjacent nodes into H . If n contains a point p , the expansion stops since every subsequent node is closer to p than to q . Figure 2 illustrates an example where the first data point (p_1) is discovered when node n_4 is de-heaped. Unlike *eager*, *lazy* does not have to retrieve the NN of n_4 . However, it still has to verify whether $p_1 \in \text{RNN}(q)$ by an expansion that visits nodes n_5 , n_2 and n_3 before determining that $q = \text{NN}(p_1)$.

Lazy takes advantage of verification queries to prune the search space. Specifically, let n be a node visited by the verification phase of a data point p . If n has not been visited by the expansion around the query, it means that $d(n,p) < d(n,q)$ since nodes are visited in ascending order of their distances and the verification query has maximum

range $d(p,q)$ (i.e., $d(n,p) < d(p,q) \leq d(n,q)$). Therefore, (by Lemma 1) n cannot lead to any RNNs of q . For instance, in Figure 2 the verification of p_1 will encounter n_5 ; thus, when n_5 is de-heaped later it will be immediately discarded. On the other hand, if n has already been visited by the expansion around q , we compare the distances $d(n,p)$ and $d(n,q)$. If $d(n,p) < d(n,q)$, all nodes that were inserted into H by the expansion of n can also be eliminated. In Figure 2, since $d(n_2, p_1) < d(n_2, q)$, node n_7 (inserted during the processing of n_2) cannot lead to a RNN. Similarly, n_6 (inserted during the processing of n_3) does not need to be expanded because $d(n_3, p_1) < d(n_3, q)$ and *lazy* terminates with $\text{RNN}(q) = \{p_1\}$.

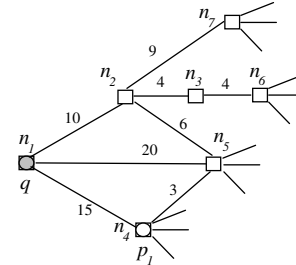


Figure 2 Example of single RNN by *lazy*

In addition, we propose two optimized versions of the basic algorithms. In particular, *eager-M* materializes the K NNs of each node n , where K is the maximum number of RNNs to be requested by any query (typically, $K \ll |P|$ where $|P|$ is the dataset cardinality). This choice supports efficient updates in the presence of object insertions and deletions. The *lazy-EP* algorithm (*EP* stands for extended pruning) expands the network in parallel using an additional heap that applies the pruning effect of the discovered points. All versions can be extended for arbitrary values of k and retrieval of bichromatic queries.

Extensive experiments with various datasets show that the problem characteristics have a significant effect on the behavior of the algorithms. *Lazy* has lower CPU overhead than *eager* in most settings (therefore, it may be preferable in the presence of large buffers), but it is very expensive for graphs (i.e., computer networks) that incur exponential expansion. *Lazy-EP* usually provides improvements over the basic algorithm. *Eager* has a more balanced behavior than both *lazy* and *lazy-EP*, but the best choice for all settings is *eager-M*, which, however, requires the materialization of the NN points of all nodes.

References

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