Fast Approximate $k$NN Graph Construction for High Dimensional Data via Recursive Lanczos Bisection

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Abstract

Nearest neighbor graphs are widely used in data mining and machine learning. A brute-force method to compute the exact $k$NN graph takes $Θ(dn^2)$ time for $n$ data points in the $d$ dimensional Euclidean space. We propose two divide and conquer methods for computing an approximate $k$NN graph in $Θ(dn^t)$ time for high dimensional data (large $d$). The exponent $t \in (1,2)$ is an increasing function of an internal parameter $α$ which governs the size of the common region in the divide step. Experiments show that a high quality graph can usually be obtained with small overlaps, i.e., for small values of $t$. A few of the practical details of the algorithms are as follows. First, the divide step uses an inexpensive Lanczos procedure to perform recursive spectral bisection. After each conquer step, an additional refinement step is performed to improve the accuracy of the graph. Finally, a hash table is used to avoid repeating distance calculations during the divide and conquer process. The combination of these techniques is shown to yield quite effective algorithms for building $k$NN graphs.

Keywords: Nearest Neighbors Graph, High Dimensional Data, Divide and Conquer, Lanczos Algorithm, Spectral Method

1. Introduction

Building nearest neighbor graphs is often a necessary step when dealing with problems arising from applications in such areas as data mining (Brito et al., 1997; Dasarathy, 2002), manifold learning (Belkin and Niyogi, 2003; Roweis and Saul, 2000; Saul and Roweis, 2003; Tenenbaum et al., 2000), robot motion planning (Choset et al., 2005), and computer graphics (Sankaranarayanan et al., 2007). Given a set of $n$ data points $X = \{x_1, \ldots, x_n\}$, a nearest neighbor graph consists of the vertex set $X$ and an edge set which is a subset of $X \times X$. The edges are defined based on a proximity measure $ρ(x_i, x_j)$ between two data points $x_i$ and $x_j$, where a small $ρ$ value means that the two points
are close. Two types of nearest neighbor graphs (Belkin and Niyogi, 2003; He and Niyogi, 2004) are often used:

1. **ε-graph**: This is an undirected graph whose edge set consists of pairs \((x_i, x_j)\) such that \(\rho(x_i, x_j)\) is less than some pre-defined threshold \(\epsilon \in \mathbb{R}^+\).

2. **kNN graph**: This is a directed graph (in general). There is an edge from \(x_i\) to \(x_j\) if and only if \(\rho(x_i, x_j)\) is among the \(k\) smallest elements of the set \(\{\rho(x_i, x_\ell) \mid \ell = 1, \ldots, i-1, i+1, \ldots, n\}\).

The \(\epsilon\)-graph is geometrically motivated, and many efficient algorithms have been proposed for computing it (see, e.g., Bentley et al., 1977; Chazelle, 1983). However, the \(\epsilon\)-graph easily results in disconnected components (Belkin and Niyogi, 2003) and it is difficult to find a good value of \(\epsilon\) which yields graphs with an appropriate number of edges. Hence, they are not suitable in many situations. On the other hand, kNN graphs have been shown to be especially useful in practice. Therefore, this paper will focus on the construction of kNN graphs, for the case when \(\rho(\cdot, \cdot)\) is the Euclidean distance between two points in the \(\mathbb{R}^d\).

When \(k = 1\), only the nearest neighbor for each data point is considered. This particular case, the *all nearest neighbors* problem, has been extensively studied in the literature. To compute the 1NN graph, Bentley (1980) proposed a multidimensional divide-and-conquer method that takes \(O(n \log^{d-1} n)\) time, Clarkson (1983) presented a randomized algorithm with expected \(O(c^d n \log n)\) time (for some constant \(c\)), and Vaidya (1989) introduced a deterministic worst-case \(O((c'd)^d n \log n)\) time algorithm (for some constant \(c'\)). These algorithms are generally adaptable to \(k > 1\). Thus, Paredes et al. (2006) presented a method to build a kNN graph, which was empirically studied and reported to require \(O(n^{1.27})\) distance calculations for low dimensional data and \(O(n^{1.90})\) calculations for high dimensional data. Meanwhile, several parallel algorithms have also been developed (Callahan, 1993; Callahan and Kosaraju, 1995; Plaku and Kavraki, 2007). Despite a rich existing literature, efficient algorithms for high dimensional data are still under-explored. In this paper we propose two methods that are especially effective in dealing with high dimensional data.

Note that the problem of constructing a kNN graph is different from the problem of *nearest neighbor* search (see, e.g., Indyk, 2004; Shakhnarovich et al., 2006, and references therein), where given a set of data points, the task is to find the \(k\) nearest points for any query point. Usually, the nearest neighbors search problem is handled by first building a data structure (such as a search tree) for the given points in a preprocessing phase. Then, queries can be answered efficiently by exploiting the search structure. Of course, the construction of a kNN graph can be viewed as a nearest neighbors search problem where each data point itself is a query. However, existing search methods in general suffer from an unfavorable trade-off between the complexity of the search structure and the accuracy of the query retrieval: either the construction of the search structure is expensive, or accurate searches are costly. The spill-tree (sp-tree) data structure (Liu et al., 2004) is shown to be empirically effective in answering queries, but since some heuristics (such as the hybrid scheme) are introduced to ensure search accuracy, the construction time cost is difficult to theoretically analyze. On the other hand, many \((1 + \epsilon)\) nearest neighbor search methods\(^1\) have been proposed with guaranteed complexity bounds (Indyk, 2004). These methods report a point within \((1 + \epsilon)\) times the distance from the query to the actual nearest neighbor. Kleinberg (1997) presented two algorithms. The first one has a \(O((d \log^2 d)(d + \log n))\) query time complexity but requires a data structure of

\(^1\) The \(\epsilon\) parameter here is different from that in the “\(\epsilon\)-graph”.  

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size $O(n \log d)^{2d}$, whereas the second algorithm uses a nearly linear (to $dn$) data structure and responds to a query in $O(n + d \log^3 n)$ time. By using locality sensitive hashing, Indyk and Motwani (1998) gave an algorithm that uses $O(n^{1+1/(1+\epsilon)} + dn)$ preprocessing time and requires $O(dn^{1/(1+\epsilon)})$ query time. Another algorithm given in Indyk and Motwani (1998) is a $O(d \text{poly log}(dn))$ search algorithm that uses $O(n^2 (n \log d \log n) / d^{1+\epsilon})$ data structure that can answer a query in $O(d \text{poly log}(dn))$ time. In general, most of the given bounds are theoretical, and have an inverse dependence on $\epsilon$, indicating expensive costs when $\epsilon$ is small. Some of these methods can potentially be applied for the purpose of efficient $k$NN graph construction, but many aspects, such as appropriate choices of the parameters that control $\epsilon$, need to be carefully considered before a practically effective algorithm is derived. Considerations of this nature will not be explored in the present paper.

The rest of the paper is organized as follows. Section 2 proposes two methods for computing approximate $k$NN graphs, and Section 3 analyzes their time complexities. Then, we show a few experiments to demonstrate the effectiveness of the methods in Section 4, and discuss two applications in Section 5. Finally, conclusions are given in Section 6.

2. Divide and Conquer $k$NN

Our general framework for computing an approximate $k$NN graph is as follows: We divide the set of data points into subsets (possibly with overlaps), recursively compute the (approximate) $k$NN graphs for the subsets, then conquer the results into a final $k$NN graph. This divide and conquer framework can clearly be separated in two distinct sub-problems: how to divide and how to conquer. The conquer step is simple: If a data point belongs to more than one subsets, then its $k$ nearest neighbors are selected from its neighbors in each subset. However, the divide step can be implemented in many different ways, resulting in different qualities of graphs. In what follows two methods are proposed which are based on a spectral bisection (Boley, 1998; Juhász and Mályusz, 1980; Tritchler et al., 2005) of the graph obtained from an inexpensive Lanczos procedure (Lanczos, 1950).

2.1 Spectral Bisection

Consider the data matrix

$$X = [x_1, \ldots, x_n] \in \mathbb{R}^{d \times n}$$

where each column $x_i$ represents a data point in $\mathbb{R}^d$. When the context is clear, we will use the same symbol $X$ to also denote the set of data points. The data matrix $X$ is centered to yield the matrix:

$$\hat{X} = [\hat{x}_1, \ldots, \hat{x}_n] = X - ce^T,$$

where $c$ is the centroid and $e$ is a column of all ones. A typical spectral bisection technique splits $\hat{X}$ into halves using a hyperplane. Let $(\sigma, u, v)$ denote the largest singular triplet of $\hat{X}$ with

$$u^T \hat{X} = \sigma v^T.$$

Then, the hyperplane is defined as $\langle u, x - c \rangle = 0$, i.e., it splits the set of data points into two subsets:

$$X_+ = \{ x_i \mid u^T \hat{x}_i \geq 0 \} \quad \text{and} \quad X_- = \{ x_i \mid u^T \hat{x}_i < 0 \}.$$
This hyperplane maximizes the sum of squared distances between the centered points \( \hat{x}_i \) to the hyperplane that passes through the centroid. This is because for any hyperplane \( \langle w, x - c \rangle = 0 \), where \( w \) is a unit vector, the squared sum is

\[
\sum_{i=1}^{n} (w^T \hat{x}_i)^2 = \|w^T \hat{X}\|_2^2 \leq \|\hat{X}\|_2^2 = \sigma^2,
\]

while \( w = u \) achieves the equality.

By a standard property of the SVD (Equation (1)), this bisection technique is equivalent to splitting the set by the following criterion:

\[
X_+ = \{ x_i \mid v_i \geq 0 \} \quad \text{and} \quad X_- = \{ x_i \mid v_i < 0 \},
\]

where \( v_i \) is the \( i \)-th entry of the right singular vector \( v \). If it is preferred that the sizes of the two subsets be balanced, an alternative is to replace the above criterion by

\[
X_+ = \{ x_i \mid v_i \geq m(v) \} \quad \text{and} \quad X_- = \{ x_i \mid v_i < m(v) \},
\]

where \( m(v) \) represents the median of the entries of \( v \).

The largest singular triplet \((\sigma, u, v)\) of \( \hat{X} \) can be computed using the Lanczos algorithm (Lanczos, 1950; Berry, 1992). In short, we first compute an orthonormal basis of the Krylov subspace \( \text{span}\{q_1, (\hat{X}^T \hat{X})q_1, \ldots, (\hat{X}^T \hat{X})^{s-1} q_1\} \) for an arbitrary initial unit vector \( q_1 \) and a small integer \( s \). Let the basis vectors form an orthogonal matrix

\[
Q_s = [q_1, \ldots, q_s].
\]

An equality resulting from this computation is

\[
Q_s^T (\hat{X}^T \hat{X}) Q_s = T_s,
\]

where \( T_s \) is a symmetric tridiagonal matrix of size \( s \times s \). Then we compute the largest eigenvalue \( \theta^{(s)} \) and corresponding eigenvector \( y_s \) of \( T_s \):

\[
T_s y_s = \theta^{(s)} y_s.
\]

Therefore, \( \theta^{(s)} \) is an approximation to the square of the largest singular value \( \sigma \), while the vector \( \tilde{v}_s = Q_s y_s \) is an approximation of the right singular vector \( v \) of \( \hat{X} \). The acute angle \( \angle(\tilde{v}_s, v) \), between \( \tilde{v}_s \) and \( v \) decays rapidly with \( s \), as is shown in an error bound established in (Saad, 1980):

\[
\sin \angle(\tilde{v}_s, v) \leq \frac{K}{C_{s-1}(1 + \gamma)}
\]

where \( K \) is a constant, \( C_k \) denotes the Chebyshev polynomial of degree \( k \) of the first kind, and \( \gamma_1 = (\bar{\lambda}_1 - \bar{\lambda}_2)/(\bar{\lambda}_2 - \bar{\lambda}_n) \) in which the \( \bar{\lambda}_i \)'s denote the eigenvalues of \( \hat{X}^T \hat{X} \), i.e., the squared singular values of \( \hat{X} \), ordered decreasingly. Note that we have \( C_{s-1}(1 + \gamma) = \cosh[(s - 1) \cosh^{-1}(1 + \gamma)] \), in which \( \cosh \) is the hyperbolic cosine. Therefore, a small value of \( s \) will generally suffice to yield an accurate approximation. Note that an exact singular vector is not needed to perform a bisection, so we usually set a fixed value, say \( s = 5 \), for this purpose. The computation of the orthonormal basis takes time \( \Theta(sd) \), while the time to compute \( \theta^{(s)} \) and \( y_s \) is negligible, since \( T_s \) is symmetric tridiagonal and \( s \) is very small. Hence the time for computing the largest singular triplet of \( \hat{X} \) is bounded by \( O(sd) \).
2.2 The Divide Step: Two Methods

Based on the above general bisection technique, we propose two ways to perform the divide step for computing an approximate \( k \)NN graph. The first, called the overlap method, divides the current set into two overlapping subsets. The second, called the glue method, divides the current set into two disjoint subsets, and uses a third set called the gluing set to help merge the two resulting disjoint \( k \)NN graphs in the conquer phase. See Figure 1. Both methods create a common region, whose size is an \( \alpha \)-portion of that of the current set, surrounding the dividing hyperplane. Details are given next.

![Figure 1: Two methods to divide a set \( X \) into subsets. The size of the common region in the middle surrounding the hyperplane is an \( \alpha \)-portion of the size of \( X \).](image)

2.2.1 THE OVERLAP METHOD

In this method, we divide the set \( X \) into two overlapping subsets \( X_1 \) and \( X_2 \):

\[
\begin{aligned}
&X_1 \cup X_2 = X, \\
&X_1 \cap X_2 \neq \emptyset.
\end{aligned}
\]  

(4)

Since \( \sigma v_i \) is the signed distance from \( \hat{x}_i \) to the hyperplane, let the set \( V \) be defined as

\[
V = \{|v_i| \mid i = 1, 2, \ldots, n\}.
\]  

(5)

Then we use the following criterion to form \( X_1 \) and \( X_2 \):

\[
X_1 = \{x_i \mid v_i \geq -h_\alpha(V)\} \quad \text{and} \quad X_2 = \{x_i \mid v_i < h_\alpha(V)\},
\]  

(6)

where \( h_\alpha(\cdot) \) is a function which returns the element that is only larger than \((100\alpha)^\%\) of the elements in the input set. The purpose of criterion (6) is to ensure that the overlap of the two subsets consists of \((100\alpha)^\%\) of all the data, i.e., that\(^2\)

\[
|X_1 \cap X_2| = \lceil \alpha |X| \rceil.
\]

---

\(^2\) Here, we assume that the distances between points and the hyperplane are all different. Ties are broken arbitrarily. The same is done for the glue method.
2.2.2 THE GLUE METHOD

In this method, we divide the set $X$ into two disjoint subsets $X_1$ and $X_2$ with a gluing subset $X_3$:

$$\begin{cases} 
X_1 \cup X_2 = X, \\
X_1 \cap X_2 = \emptyset, \\
X_1 \cap X_3 \neq \emptyset, \\
X_2 \cap X_3 \neq \emptyset.
\end{cases} \quad (7)$$

The criterion to build these subsets is as follows:

$$X_1 = \{x_i \mid v_i \geq 0\}, \quad X_2 = \{x_i \mid v_i < 0\}, \quad X_3 = \{x_i \mid -h_\alpha(V) \leq v_i < h_\alpha(V)\}. \quad (8)$$

Note that the gluing subset $X_3$ in this method is exactly the intersection of the two subsets in the overlap method. Hence, $X_3$ also contains $(100\alpha)\%$ of the data.

2.3 Refinement

In order to improve the quality of the resulting graph, during each recursion after the conquer step, the graph can be refined at a small cost. The idea is to update the $k$ nearest neighbors for each point by selecting from a pool consisting of its neighbors and the neighbors of its neighbors. Formally, if $N(x)$ is the set of current nearest neighbors of $x$ before refinement, then, for each point $x$, we re-select its $k$ nearest neighbors from

$$N(x) \cup \left( \bigcup_{z \in N(x)} N(z) \right).$$

2.4 The Algorithms

We are ready to present the complete algorithms for both methods; see Algorithms 1 and 2. These algorithms share many similarities: They both fall in the framework of divide and conquer; they both call the brute-force procedure $k$NN-BRUTEFORCE to compute the graph when the size of the set is smaller than a threshold ($n_k$); they both recursively call themselves on smaller subsets; and they both employ a CONQUER procedure to merge the graphs computed for the subsets and a REFINE procedure to refine the graph during each recursion. The difference is that Algorithm 1 calls DIVIDE-OVERLAP to divide the set into two subsets (Section 2.2.1), while Algorithm 2 calls DIVIDE-GLUE to divide the set into three subsets (Section 2.2.2). For the sake of completeness, pseudocodes of all the mentioned procedures are given in Appendix A.

2.5 Storing Computed Distances in a Hash Table

The brute-force method computes $\Theta(n^2)$ pairs of distances, each of which takes $\Theta(d)$ time. One advantage of the methods presented in this paper over brute-force methods is that the distance calculations can be significantly reduced thanks to the divide and conquer approach. The distances are needed/computed in: (1) the $k$NN-BRUTEFORCE procedure which computes all the pairwise distances and selects the $k$ smallest ones for each point, (2) the CONQUER procedure which selects
Algorithm 1 Approximate kNN Graph Construction: The Overlap Method

1: function $G = k\text{NN-OVERLAP}(X, k, \alpha)$
2: if $|X| < n_k$ then
3:   $G \leftarrow k\text{NN-BRUTEFORCE}(X, k)$
4: else
5:   $(X_1, X_2) \leftarrow \text{DIVIDE-OVERLAP}(X, \alpha)$ \hspace{1cm} \triangleright Section 2.2.1
6:   $G_1 \leftarrow k\text{NN-OVERLAP}(X_1, k, \alpha)$
7:   $G_2 \leftarrow k\text{NN-OVERLAP}(X_2, k, \alpha)$
8:   $G \leftarrow \text{CONQUER}(G_1, G_2, k)$ \hspace{1cm} \triangleright Section 2, beginning
9:   $\text{REFINE}(G, k)$ \hspace{1cm} \triangleright Section 2.3
10: end if
11: end function

Algorithm 2 Approximate kNN Graph Construction: The Glue Method

1: function $G = k\text{NN-GLUE}(X, k, \alpha)$
2: if $|X| < n_k$ then
3:   $G \leftarrow k\text{NN-BRUTEFORCE}(X, k)$
4: else
5:   $(X_1, X_2, X_3) \leftarrow \text{DIVIDE-GLUE}(X, \alpha)$ \hspace{1cm} \triangleright Section 2.2.2
6:   $G_1 \leftarrow k\text{NN-GLUE}(X_1, k, \alpha)$
7:   $G_2 \leftarrow k\text{NN-GLUE}(X_2, k, \alpha)$
8:   $G_3 \leftarrow k\text{NN-GLUE}(X_3, k, \alpha)$
9:   $G \leftarrow \text{CONQUER}(G_1, G_2, G_3, k)$ \hspace{1cm} \triangleright Section 2, beginning
10: $\text{REFINE}(G, k)$ \hspace{1cm} \triangleright Section 2.3
11: end if
12: end function

$k$ smallest distances from at most $2k$ candidates for each point, and (3) the $\text{REFINE}$ procedure which selects the $k$ smallest distances from at most $k + k^2$ candidates for each point. Many of the distances computed from $k\text{NN-BRUTEFORCE}$ and $\text{REFINE}$ are reused in $\text{CONQUER}$ and $\text{REFINE}$, with probably more than once for some pairs. A naive way is to allocate memory for an $n \times n$ matrix that stores all the computed distances to avoid duplicate calculations. However this consumes too much memory and is not necessary. A better approach is to use a hash table to store the computed distances. This will save a significant amount of memory, as a later experiment shows that only a small portion of the $n^2$ pairs are actually computed. Furthermore the computational time will not be affected since hash tables are efficient for both retrieving wanted items from and inserting new items into the table (we do not need the delete operation).

The ideal hash function maps (the distance between) a pair of data points $(x_i, x_j)$ to a bucket such that the probability of collision is low. For simplicity of implementations, we use a hash function that maps the key $(x_i, x_j)$ to $i$ (and at the same time to $j$). Collisions easily occur since many distances between the point $x_i$ and other points are computed during the whole process. However, this naive hashing has already shown rather appealing results in run time. More elaborate implementations should consider more effective hashing (e.g. double hashing) schemes.
3. Complexity Analysis

A thorough analysis shows that the time complexities for the overlap method and the glue method are sub-quadratic (in \( n \)), and the glue method is always asymptotically faster than the overlap method. To this end we assume that in each divide step the subsets \( X_1 \) and \( X_2 \) (in both methods) are balanced. This assumption can always be satisfied by using the Equation (3) to bisect the data set, instead of the criterion (2b). Hence, the time complexity \( T_o \) for the overlap method and \( T_g \) for the glue method satisfy the following recurrence relations:

\[
T_o(n) = 2T_o((1 + \alpha)n/2) + f(n), \quad (9)
\]

\[
T_g(n) = 2T_g(n/2) + T_g(\alpha n) + f(n), \quad (10)
\]

where \( f(n) \) is the combined time for the divide, conquer, and refine steps.

3.1 The Complexity of \( f \)

The function \( f(n) \) consists of the following three components.

(a) The time for the divide step. This includes the time to compute the largest right singular vector \( v \) of the centered matrix \( \hat{X} \), and the time to divide points into subsets \( X_1 \) and \( X_2 \) (in the overlap method) or subsets \( X_1 \), \( X_2 \) and \( X_3 \) (in the glue method). The former has been shown to be \( O(sd\nu) \) in Section 2.1, where the number of Lanczos steps \( s = 5 \) is fixed in the implementation, while for the latter we can use a linear time selection method to find the value \( h_{\alpha}(V) \). Therefore the overall time for this step is \( O(\alpha n) \).

(b) The time for the conquer step. This step only involves the points in \( X_1 \setminus X_2 \) in the overlap method or \( X_3 \) in the glue method. For each of the \( \alpha n \) points in these subsets, \( k \) nearest neighbors are chosen from at most \( 2k \) candidates. Therefore the time is \( O(k\alpha n) \).

(c) The time for the refine step. For each point, \( k \) nearest neighbors are chosen from at most \( k + k^2 \) candidates. If all these distances need to be computed, the overall time is \( O(k^2dn) \). To the other extreme, if none of them are computed, the factor \( d \) can be omitted, which results in \( O(k^2n) \). In practice, by using a hash table, only a very small fraction of the \( k + k^2 \) distances are actually computed in this step; see also Table 3. Hence, the best cost estimate for this step is \( O(k^2n) \).

Indeed, the ultimate useful information from this estimate is that the time for the refinement is much less than that for the division (which has a \( O(d\nu n) \) cost), or at best is of the same order as the latter. This can be seen from another perspective: Let the number of neighbors \( k \) be a constant. Then even if all the distances are computed, the time is \( O(k^2dn) = O(d\nu n) \).

From the above three components and the fact that the dimension \( d \) dominates \( k \), we conclude that \( f(n) \) is bounded by \( O(d\nu n) \).

3.2 The Complexities of \( T_o \) and \( T_g \)

By substituting \( f(n) = O(d\nu n) \) into (9) and (10), we derive the closed form solutions to \( T_o(n) \) and \( T_g(n) \), which are stated in the following two theorems.

**Theorem 1** The time complexity for the overlap method is

\[
T_o(n) = \Theta(d\nu n), \quad (11)
\]
where
\[ t_0 = \log_2/(1+\alpha)^2 = \frac{1}{1 - \log_2(1 + \alpha)}. \] (12)

**Theorem 2** The time complexity for the glue method is
\[ T_g(n) = \Theta(dn^\alpha), \] (13)
where \( t_g \) is the solution to the equation
\[ \frac{2}{2t} + \alpha' = 1. \] (14)

The proofs will be given in Appendix B. Figure 2 plots the two curves of \( t_0 \) and \( t_g \) as functions of \( \alpha \), together with a table that lists some of their values. Figure 2 suggests that the glue method is asymptotically faster than the overlap method. Indeed, this is true for any choice of \( \alpha \).

![Figure 2: The exponents \( t_0 \) and \( t_g \) as functions of \( \alpha \).](image)

<table>
<thead>
<tr>
<th>( \alpha )</th>
<th>0.05</th>
<th>0.10</th>
<th>0.15</th>
<th>0.20</th>
<th>0.25</th>
<th>0.30</th>
</tr>
</thead>
<tbody>
<tr>
<td>( t_0 )</td>
<td>1.08</td>
<td>1.16</td>
<td>1.25</td>
<td>1.36</td>
<td>1.47</td>
<td>1.61</td>
</tr>
<tr>
<td>( t_g )</td>
<td>1.06</td>
<td>1.12</td>
<td>1.17</td>
<td>1.22</td>
<td>1.27</td>
<td>1.33</td>
</tr>
</tbody>
</table>

**Theorem 3** When \( 0 < \alpha < 1 \), the exponents \( t_0 \) in Theorem 1 and \( t_g \) in Theorem 2 obey the following relation:
\[ 1 < t_g < t_0. \] (15)

The proof of the above theorem will be given in Appendix B. We remark that when \( \alpha > \sqrt{2} - 1 \approx 0.41, t_0 > 2 \). In this situation the overlap method becomes asymptotically slower than the brute-force method. Similarly, when \( \alpha > 1/\sqrt{2} \approx 0.71, t_g > 2 \). Hence, a large \( \alpha \) (> 0.41) is not useful in practice.

4. Experiments

In this section we show a few experimental results to illustrate the running times of the two proposed methods compared with the brute-force method, and the qualities of the resulting graphs. The experiments were performed under a Linux workstation with two P4 3.20GHz CPUs and 2GB memory. The algorithms were all implemented using C/C++, and the programs were compiled using g++ with -O2 level optimization. The divide and conquer methods were implemented according to Algorithms 1 and 2, and the brute-force method was exactly as in the procedure \texttt{kNN-BRUTEFORCE} given in Appendix A.
4.1 Running Time

Figure 3 plots the running times versus the dimension $d$ and the number of data points $n$ on a synthetic data set. Since the distribution of the data should have little impact on the running times of the methods, we used random data (drawn from the uniform distribution over $[0, 1]^d$) for this experiment. From Figure 3(a), it is clear that the running time is linear with respect to the dimension $d$. This is expected from the complexity analysis. In Figure 3(b), we used $\alpha = 0.2$, which corresponds to the theoretical values $t_0 = 1.36$ and $t_0 = 1.22$. We used curves in the form $c_1n^{1.36} + c_2n + c_3$ and $c_4n^{1.22} + c_5n + c_6$ to fit the running times of the overlap method and the glue method, respectively. The fitted curves are also shown in Figure 3. It can be seen that the experimental results match the theoretical analysis quite well.

![Figure 3: The running times for randomly generated data.](image)

4.2 Quality

In another experiment, we used four real-life data sets to test the qualities of the resulting $k$NN graphs: The FREY$^3$ face video frames (Roweis and Saul, 2000), the extYaleB$^4$ database (Lee et al., 2005), the MNIST$^5$ digit images (LeCun et al., 1998), and the PIE$^6$ face database (Sim et al., 2003). These data sets are all image sets that are widely used in the literature in the areas of face recognition, dimensionality reduction, etc. For MNIST, we used only the test set. Table 1 gives some characteristics of the data. The number of images is equal to $n$ and the size of each image, i.e., the total number of pixels for each image, is the dimension $d$. The dimensions vary from about 500 to 10,000. Since some of these data sets were also used later to illustrate the practical usefulness of our methods in real applications, for each one we used a specific $k$ value that was used in the experiments of past publications. These values are typically around 10.

The qualities of the resulting graphs versus the running times are plotted in Figure 4. Each plotted point in the figure corresponds to a choice of $\alpha$ ($= 0.05, 0.10, 0.15, 0.20, 0.25, 0.30$). The running times of the brute-force method are also indicated. We use two quality criteria: accuracy and average rank. The accuracy of an approximate $k$NN graph $G'$ (with regard to the exact graph $G$) is defined as

$$\text{accuracy}(G') = \frac{|E(G') \cap E(G)|}{|E(G)|},$$

where $E(\cdot)$ means the set of directed edges in the graph. Thus, the accuracy is within the range $[0, 1]$, and higher accuracy means better quality. The rank $r_u(v)$ of a vertex $v$ with regard to a vertex $u$ is the position of $v$ in the vertex list sorted in ascending order of the distance to $u$. (By default, the rank of the nearest neighbor of $u$ is 1.) Thus, the average rank is defined as

$$\text{ave-rank}(G') = \frac{1}{kn} \sum_u \sum_{v \in N(u)} r_u(v),$$

where $N(u)$ means the neighborhood of $u$ in the graph $G'$. The exact $k$NN graph has the average rank $(1 + k)/2$.

It can be seen from Figure 4 that the qualities of the resulting graphs exhibit similar trends by using both measures. Take the graph accuracy for example. The larger $\alpha$, the more accurate the resulting graph. However, larger values of $\alpha$ lead to more time-consuming runs. In addition, the glue method is much faster than the overlap method for the same $\alpha$, while the latter yields more accurate graphs than the former. The two methods are both significantly faster than the brute-force method when an appropriate $\alpha$ is chosen, and they can yield high quality graphs even when $\alpha$ is small.

It is interesting to note that in all the plots, the red circle-curve and the blue plus-curve roughly overlap. This similar quality-time trade-off seems to suggest that neither of the method is superior to the other one; the only difference is the $\alpha$ value used to achieve the desired quality. However, we note that different approximate graphs can yield the same accuracy value or average rank value, hence the actual quality of the graph depends on real applications.

### 4.3 Distance Calculations and Refinements

We also report the percentage of the distance calculations, which is one of the dominant costs of the graph construction. The superior efficiency of our methods lies in two facts. First, for $n$ data points, the brute-force method must compute distances between $n(n - 1)/2$ pairs of points, while our methods compute only a small fraction of this number. Second, the use of a (hash) table brings the benefit of avoiding repeating distance calculations if the evaluations of the same distance are required multiple times. Tables 2 and 3 confirm these two claims. As expected, the larger the common region ($\alpha$) is, the more distances need to be calculated, while the benefit of the hashing

<table>
<thead>
<tr>
<th></th>
<th>FREY</th>
<th>extYaleB</th>
<th>MNIST</th>
<th>PIE</th>
</tr>
</thead>
<tbody>
<tr>
<td># imgs ($n$)</td>
<td>1,965</td>
<td>2,414</td>
<td>10,000</td>
<td>11,554</td>
</tr>
<tr>
<td>img size ($d$)</td>
<td>20 $\times$ 28</td>
<td>84 $\times$ 96</td>
<td>28 $\times$ 28</td>
<td>32 $\times$ 32</td>
</tr>
</tbody>
</table>

Table 1: Image data sets.
Figure 4: Graph quality versus running time for different data sets. Each row of the plots corresponds to one data set. The left column of plots shows the $1-\text{accuracy}$ measure. The right column shows the average-rank measure. Each plotted point corresponds to a choice of $\alpha$. From left to right on each curve, the $\alpha$ values are 0.05, 0.10, 0.15, 0.20, 0.25, 0.30, respectively.
becomes more significant. It can also be seen that the savings in distance calculations are more remarkable when \( n \) becomes larger and \( k \) becomes smaller, e.g., \( k = 5 \) for the data set PIE.

<table>
<thead>
<tr>
<th>( \alpha )</th>
<th>FREY ((k = 12))</th>
<th>extYaleB ((k = 10))</th>
<th>MNIST ((k = 8))</th>
<th>PIE ((k = 5))</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.05</td>
<td>6.07%</td>
<td>5.10%</td>
<td>5.13%</td>
<td>4.42%</td>
</tr>
<tr>
<td>0.10</td>
<td>6.55%</td>
<td>5.80%</td>
<td>5.58%</td>
<td>5.08%</td>
</tr>
<tr>
<td>0.15</td>
<td>7.48%</td>
<td>6.35%</td>
<td>6.05%</td>
<td>5.46%</td>
</tr>
<tr>
<td>0.20</td>
<td>8.28%</td>
<td>6.57%</td>
<td>6.66%</td>
<td>5.66%</td>
</tr>
<tr>
<td>0.25</td>
<td>9.69%</td>
<td>7.00%</td>
<td>7.36%</td>
<td>6.02%</td>
</tr>
<tr>
<td>0.30</td>
<td>11.48%</td>
<td>7.46%</td>
<td>8.34%</td>
<td>6.26%</td>
</tr>
</tbody>
</table>

Table 2: Percentages of distance calculations (with respect to \( n(n - 1)/2 \), for different data sets, different methods, and different \( \alpha \)'s.

<table>
<thead>
<tr>
<th>( \alpha )</th>
<th>FREY ((k = 12))</th>
<th>extYaleB ((k = 10))</th>
<th>MNIST ((k = 8))</th>
<th>PIE ((k = 5))</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.05</td>
<td>9.44%</td>
<td>12.15%</td>
<td>11.46%</td>
<td>14.80%</td>
</tr>
<tr>
<td>0.10</td>
<td>7.07%</td>
<td>12.09%</td>
<td>8.46%</td>
<td>14.36%</td>
</tr>
<tr>
<td>0.05</td>
<td>5.18%</td>
<td>10.81%</td>
<td>6.25%</td>
<td>12.79%</td>
</tr>
<tr>
<td>0.20</td>
<td>3.65%</td>
<td>9.12%</td>
<td>4.35%</td>
<td>11.04%</td>
</tr>
<tr>
<td>0.05</td>
<td>2.62%</td>
<td>7.52%</td>
<td>2.90%</td>
<td>9.55%</td>
</tr>
<tr>
<td>0.30</td>
<td>1.83%</td>
<td>6.44%</td>
<td>1.88%</td>
<td>7.89%</td>
</tr>
</tbody>
</table>

Table 3: Percentages of actual distance calculations with respect to the total number of needed distances in the refine step, for different data sets, different methods, and different \( \alpha \)'s. This is equivalent to the failure rate of hash table lookups.

The final experiment illustrates the importance of the refine step. Figure 5 shows the decrease in quality if the \textsc{Refine} procedure is not invoked. The refinement greatly improves the accuracy of the approximate graph (especially for \( \alpha \) less than 0.2) at some additional cost in the execution time. This additional expense is worthwhile if the goal is to compute a high quality \( k \)NN graph.

5. Applications

\( k \)NN graphs have been widely used in various data mining and machine learning applications. This section discusses two scenarios where the approximate \( k \)NN graphs resulting from our proposed techniques can provide an effective replacement for the exact \( k \)NN graph.

5.1 Agglomerative Clustering

Agglomerative clustering (Ward, 1963) is a clustering method which exploits a hierarchy for a set of data points. Initially each point belongs to a separate cluster. Then, the algorithm iteratively merges a pair of clusters that has the lowest merge cost, and updates the merge costs between the
newly merged cluster and the rest of the clusters. This process terminates when the desired number of clusters is reached (or when all the points belong to the single final cluster). A straightforward implementation of the method takes $O(n^3)$ time, since there are $O(n)$ iterations, each of which takes $O(n^2)$ time to find the pair with the lowest merge cost (Shanbehzadeh and Ogunbona, 1997). Fränti et al. (2006) proposed, at each iteration, to maintain the $k$NN graph of the clusters, and merge two clusters that are the nearest neighbors in the graph. Let the number of clusters at the present iteration be $m$. Then this method takes $O(km)$ time to find the closest clusters, compared with the $O(m^2)$ cost of finding the pair with the lowest merge cost. With a delicate implementation using a doubly linked list, they showed that the overall running time of the clustering process reduces to $O(\tau n \log n)$, where $\tau$ is the number of nearest neighbor updates at each iteration. Their method greatly speeds up the clustering process, while the clustering quality is not much degraded.

However, the quadratic time to create the initial $k$NN graph eclipses the improvement in the clustering time. One solution is to use an approximate $k$NN graph that can be inexpensively created. Virmajoki and Fränti (2004) proposed a divide-and-conquer algorithm to create an approximate $k$NN graph, but their time complexity was overestimated. Our approach also follows the common framework of divide and conquer. However, we bring three improvements over previous work: (1) two methods to perform the divide step are proposed, (2) an efficient way to compute the separating hyperplane is described, and (3) a detailed and rigorous analysis on the time complexity is provided. This analysis in particular makes the proposed algorithms practical, especially in the presence of high dimensional data (e.g., when $d$ is in the order of hundreds or thousands).

We performed an experiment on the data set PIE with 68 classes (see Figure 6). Since class labels are known, we used the purity and the entropy (Zhao and Karypis, 2004) as quality measures. They are defined as

$$Purity = \frac{1}{n} \sum_{i=1}^{q} n_i \text{Purity}(i), \quad \text{where} \quad \text{Purity}(i) = \frac{1}{n_i} \max_{j} \left(n_{ij}^f\right),$$

Figure 5: The refine step boosts the accuracy of the graph at some additional computational cost. Data set: MNIST. The settings are the same as in Figure 4.
and

\[
\text{Entropy} = \sum_{i=1}^{q} \frac{n_i}{n} \text{Entropy}(i), \quad \text{where} \quad \text{Entropy}(i) = -\sum_{j=1}^{q} \frac{n_{ij}}{n_i} \log_q \frac{n_{ij}}{n_i}.
\]

Here, \( q \) is the number of classes/clusters, \( n_i \) is the size of cluster \( i \), and \( n_{ij} \) is the number of class \( j \) data that are assigned to the \( i \)-th cluster. The purity and the entropy both range from 0 to 1. In Figure 6 we show the purities, the entropies, and the values of the objective function for a general purpose clustering (sum of squared errors, SSE), for different methods and different \( \alpha \)'s. In general, a higher purity, a lower entropy, and/or a lower SSE means a better clustering quality. As can be seen the qualities of the clusterings obtained from the approximate \( k \)NN graphs are very close to those resulting from the exact graph, with a few being even much better. It is interesting to note that
the clustering results seem to have little correlation with the qualities of the graphs governed by the value \( \alpha \).

### 5.2 Dimensionality Reduction

Many dimensionality reduction methods, e.g., locally linear embedding (LLE) (Roweis and Saul, 2000), Laplacian eigenmaps (Belkin and Niyogi, 2003), locality preserving projections (LPP) (He and Niyogi, 2004), and orthogonal neighborhood preserving projections (ONPP) (Kokiopoulou and Saad, 2007), compute a low dimensional embedding of the data by preserving the local neighborhoods for each point. For example, in LLE, a weighted adjacency matrix \( W \) is first computed to minimize the following objective:

\[
E(W) = \sum_i \left\| x_i - \sum_{j \in N(x_i)} W_{ij} x_j \right\|^2, \quad \text{subject to } \sum_j W_{ij} = 1, \forall i,
\]

where \( N(\cdot) \) means the neighborhood of a point. Then, a low dimensional embedding \( Y = [y_1, \ldots, y_n] \) is computed such that it minimizes the objective:

\[
\Phi(Y) = \sum_i \left\| y_i - \sum_{j \in N(y_i)} W_{ij} y_j \right\|^2, \quad \text{subject to } YY^T = I.
\]

The final solution, \( Y \), is the matrix whose column-vectors are the \( r \) bottom right singular vectors of the Laplacian matrix \( L_{\text{LLE}} = I - W \). As another example, in Laplacian eigenmaps, the low dimensional embedding \( Y = [y_1, \ldots, y_n] \) is computed so as to minimize the cost function:

\[
\Psi(Y) = \sum_{i,j} W_{ij} \left\| y_i - y_j \right\|^2, \quad \text{subject to } YY^T = I,
\]

where \( D \) is the diagonal degree matrix. Here, the weighted adjacency matrix \( W \) is defined in a number of ways, one of the most popular being the weights of the heat kernel

\[
W_{ij} = \begin{cases} 
\exp\left(-\frac{\|x_i - x_j\|^2}{\sigma^2}\right) & \text{if } x_i \in N(x_j) \text{ or } x_j \in N(x_i), \\
0 & \text{otherwise}.
\end{cases}
\]

The solution, \( Y \), is simply the matrix of \( r \) bottom eigenvectors of the normalized Laplacian matrix \( L_{\text{eigenmaps}} = I - D^{-1/2}WD^{-1/2} \) (subject to scaling).

A thorn in the nicely motivated formulations for the above approaches, is that they all begin with a rather expensive computation to obtain the neighborhood graph of the data. On the other hand, the cost of computing the solution \( Y \) via, e.g., the Lanczos algorithm, is relatively inexpensive, and can be summarized as \( O(rkn) \), which is independent of the dimension \( d \). The methods discussed in this paper are suitable alternatives to the expensive brute-force approach to extract the exact \( k \)NN graph, since the approximate graphs are accurate enough for the purpose of dimensionality reduction, while the time costs are significantly smaller. Figures 7 and 8 provide two illustrations of this.

---

1. To be more exact, the dominant cost of computing \( r \) singular vectors/eigenvectors of a sparse matrix by the Lanczos method is \( O(r' \cdot \text{nnz}) \), where \( r' \) is the number of Lanczos steps and \( \text{nnz} \) is the number of nonzeros in the matrix. The value \( r' \) is in practice a few times of \( r \), and in our situation, the matrix is the (normalized) graph Laplacian, hence \( \text{nnz} = O(kn) \).
In Figure 7 are the plots of the dimensionality reduction results of LLE applied to the data set FREY, where we used $k = 12$ as that in Roweis and Saul (2000, Figure 3). Figure 7(a) shows the result when using the exact $k$NN graph, while Figure 7(b) shows the result when using the approximate $k$NN graph by the overlap method with $\alpha = 0.30$. It is clear that the two results are almost identical. Figures 7(c) and 7(d) give two plots when the glue method is used. Although the embeddings are different from those of the exact $k$NN graph in Figure 7(a), they also represent the original data manifold quite well. This can be seen by tracking the relative locations of the sequence of images (as shown in 7(e)) in the two dimensional space.

Figure 7: Dimensionality reduction on the data set FREY by LLE.
Figure 8 shows the plots of the dimensionality reduction results of Laplacian eigenmaps applied to the data set MNIST, where we used $k = 5$. Figure 8(a) shows the original result by using the exact $k$NN graph, while 8(b), 8(c) and 8(d) show the results by using the overlap method with $\alpha = 0.30$, the glue method with $\alpha = 0.15$, and the glue method with $\alpha = 0.30$, respectively. The four plots all show clear clusterings of the ten classes (digits from 0 to 9), and the localization patterns of the clusterings are very similar.

6. Conclusions
We have proposed two sub-quadratic time methods under the framework of divide and conquer for computing approximate $k$NN graphs for high dimensional data. The running times of the methods, as well as the qualities of the resulting graphs, depend on an internal parameter that controls the...
overlap size of the subsets in the divide step. Experiments show that in order to obtain a high quality graph, a small overlap size is usually sufficient and this leads to a small exponent in the time complexity. An avenue of future research is to theoretically analyze the quality of the resulting graphs in relation to the overlap size. The resulting approximate graphs have a wide range of applications as they can be safely used as alternatives to the exact $k$NN graph. We have shown two such examples: one in agglomerative clustering and the other in dimensionality reduction. Thus, replacing the exact $k$NN graph construction with one produced by the methods proposed here, can significantly alleviate what currently constitutes a major bottleneck in these applications.

Acknowledgments

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Appendix A. Pseudocodes of Procedures

1: function $G = k$NN-BRUTEFORCE($X$, $k$)
2: for $i \leftarrow 1, \ldots, n$ do
3: for $j \leftarrow i + 1, \ldots, n$ do
4: Compute $\rho(x_i, x_j) = \| x_i - x_j \|$  \hspace{1cm} \rho(x_j, x_i) = \rho(x_i, x_j)$
5: end for
6: Set $N(x_i) = \{ x_j \mid \rho(x_i, x_j) \text{ is among the } k \text{ smallest elements for all } j \neq i \}$
7: end for
8: end function

1: function ($X_1, X_2$) = DIVIDE-OVERLAP($X$, $\alpha$)
2: Compute the largest right singular vector $v$ of $\hat{X} = X - ce^T$
3: Let $V = \{ |v_i| \mid i = 1, 2, \ldots, n \}$
4: Find $h_{\alpha}(V)$  \hspace{1cm} \triangleright$\text{See Section 2.2.1 for the definition}$
5: Set $X_1 = \{ x_i \mid v_i \geq -h_{\alpha}(V) \}$
6: Set $X_2 = \{ x_i \mid v_i < h_{\alpha}(V) \}$
7: end function

1: function ($X_1, X_2, X_3$) = DIVIDE-GLUE($X$, $\alpha$)
2: Compute the largest right singular vector $v$ of $\hat{X} = X - ce^T$
3: Let $V = \{ |v_i| \mid i = 1, 2, \ldots, n \}$
4: Find $h_{\alpha}(V)$
5: Set $X_1 = \{ x_i \mid v_i \geq 0 \}$
6: Set $X_2 = \{ x_i \mid v_i < 0 \}$
7: Set $X_3 = \{ x_i \mid -h_{\alpha}(V) \leq v_i < h_{\alpha}(V) \}$
8: end function

1: function $G = CONQUER(G_1, G_2, k)$
2: $G = G_1 \cup G_2$
Appendix B. Proofs

Theorem 1 follows from the Master Theorem (Cormen et al., 2001, Chapter 4.3). Theorem 2 is an immediate consequence of the following lemma, which is straightforward to verify.

**Lemma 4** The recurrence relation

\[ T(n) = 2T(n/2) + T(\alpha n) + n \]

with \( T(1) = 1 \) has a solution

\[ T(n) = \left( 1 + \frac{1}{\alpha} \right) n' - \frac{n}{\alpha} \]

where \( t \) is the solution to the equation

\[ \frac{2}{2^t} + \alpha' = 1. \]

The proof of Theorem 3 requires two lemmas.

**Lemma 5** When \( 0 < x < 1 \),

\[ \log_2(1-x^2) > \left( \log_2(1-x) \right) \left( \log_2(1+x) \right). \]

**Proof** By Taylor expansion,

\[
\left( \ln(1-x) \right) \left( \ln(1+x) \right) \\
= \left( - \sum_{n=1}^{\infty} \frac{x^n}{n} \right) \left( \sum_{n=1}^{\infty} \frac{(-1)^{n+1}x^n}{n} \right) \\
= - \sum_{n=1}^{\infty} \left( \sum_{m=1}^{2n-1} \frac{(-1)^{m-1}}{m(2n-m)} \right) x^{2n}
\]
\[\sum_{n=1}^{\infty} \left( \frac{1}{2n} \left( \frac{1}{1} + \frac{1}{2n-1} \right) - \frac{1}{2n} \left( \frac{1}{2} + \frac{1}{2n-2} \right) + \cdots + \frac{(-1)^{n-1}}{n^2} \right) x^{2n}\]

The inequality in the lemma follows by changing the bases of the logarithms.

**Lemma 6** The following inequality

\[\log_2(ab) > (\log_2 a)(\log_2 b)\]

holds whenever \(0 < a < 1 < b < 2\) and \(a + b \geq 2\).

**Proof** By using \(b \geq 2 - a\), we have the following two inequalities

\[\log_2(ab) \geq \log_2(a(2-a)) = \log_2(1 - (1-a))(1 + (1-a)) = \log_2(1 - (1-a)^2)\]

and

\[(\log_2 a)(\log_2 b) \leq (\log_2 a)(\log_2(2-a)) = \log_2(1 - (1-a)) \times \log_2(1 + (1-a)).\]

Then by applying Lemma 5 with \(1 - a = x\), we have

\[\log_2(1 - (1-a)^2) > \log_2(1 - (1-a)) \times \log_2(1 + (1-a)).\]

Thus, the inequality of the lemma holds.

**Proof of Theorem 3** From Theorem 2 we have

\[t_g = 1 - \log_2(1 - \alpha^t) > 1.\]

Then

\[t_0 - t_g = \frac{1}{1 - \log_2(1 + \alpha)} - 1 + \log_2(1 - \alpha^t)\]

\[= \frac{\log_2(1 + \alpha) + \log_2(1 - \alpha^t) - \log_2(1 + \alpha) \times \log_2(1 - \alpha^t)}{1 - \log_2(1 + \alpha)}.\]

Since \(0 < \alpha < 1\), the denominator \(1 - \log_2(1 + \alpha)\) is positive. By Lemma 6 the numerator is also positive. Hence \(t_0 > t_g\).
References


