EMPIRICAL COMPLEXITY OF COMPARATOR-BASED NEAREST NEIGHBOR DESCENT

JACOB D. BARON     R.W.R. DARLING

National Security Agency, Fort George G. Meade, MD 20755-6844, USA

Abstract. A Java parallel streams implementation of the $K$-nearest neighbor descent algorithm is presented using a natural statistical termination criterion. Input data consist of a set $S$ of $n$ objects of type $V$, and a $\text{Function}<V, \text{Comparator}<V>>$, which enables any $x \in S$ to decide which of $y, z \in S \setminus \{x\}$ is more similar to $x$. Experiments with the Kullback-Leibler divergence Comparator support the prediction that the number of rounds of $K$-nearest neighbor updates need not exceed twice the diameter of the undirected version of a random regular out-degree $K$ digraph on $n$ vertices. Overall complexity was $O(nK^2 \log K(n))$ in the class of examples studied. When objects are sampled uniformly from a $d$-dimensional simplex, accuracy of the $K$-nearest neighbor approximation is high up to $d = 20$, but declines in higher dimensions, as theory would predict.

Keywords: similarity search, nearest neighbor, ranking system, triplet comparison, comparator, random graph, proximity graph, expander graph

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1. Introduction

1.1. Context. Baron and Darling [3] provided a theoretical analysis of the $K$-nearest neighbor descent ($K$-NN Descent) algorithm for $K$-nearest neighbor approximation proposed and implemented by Dong, Charikar, and Li [7].

This sequel reports on a generic Java parallel streams implementation of $K$-NN Descent, which was written to support a forthcoming implementation of the partitioned nearest neighbors local depth algorithm [4]. While testing this implementation, we acquired statistical data which shed light on the performance of $K$-NN Descent, under a new termination criterion. This brief report does not attempt comparison of $K$-NN Descent with other algorithms, as

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Table 1. Comparison among four K-NN Descent implementations.

| Authors           | Language | Asymmetric? | Parallelization?          | Termination               |
|-------------------|----------|------------|---------------------------|---------------------------|
| Dong et al [7]    | C++      | no         | OpenMP & map-reduce       | δ-proportion update       |
| McInnes [14]      | Python   | no         | none                      | no possible update        |
| Kluser et al [12] | C        | no         | none                      | no possible update        |
| This paper        | Java     | yes        | fork-join pool            | statistical criterion     |

is reported in [7]. Nor shall we outline the different approaches to K-nearest neighbor approximation, briefly surveyed by Aumüller, Bernhardsson and, Faithfull [1]. We mention the recent competition [13] to surpass the industry leader FAISS [11], in the case of a billion dense vectors in dimension 96 to 256, under the $\ell_2$ norm.

1.2. Previous implementations. A high level summary of previous K-NN Descent implementations is shown in Table 1, along with our own in the final row.

1.2.1. Original implementation for metrics. A sophisticated OpenMP and map-reduce implementation of K-NN Descent is described by Dong et al [7]. These authors employ optimizations applicable to symmetric similarity functions, and employ two stopping criteria, both of which are different to ours. The authors describe in detail the application to five well-studied data sets of sizes between 28,755 and 857,820 using several symmetric similarity measures, and compare K-NN Descent with Recursive Lanczos Bisection and Locality Sensitive Hashing.

1.2.2. Python implementation. McInnes created the widely-used pynndescent Python version [14], which is used in UMAP [15]. It assumes that similarity is obtained from a metric, of which 22 examples are available in the code. This is one of 19 single-threaded approximate K-NN Python algorithms among benchmarks at [5].

1.2.3. Single-threaded C Implementation. Kluser et al [12] describe a runtime-optimized C implementation for the $\ell_2$-distance metric, and report performance improvements over the two versions above. Their approach increases locality by improving the otherwise irregular memory access pattern.

1.3. Novelty of our implementation. Here are some more details, beyond Table 1, of what distinguishes our implementation from the others.

(a) **Non-metric:** Input data consist of a set $S$ of $n$ objects of type $V$, and a Function<$V$, Comparator<$V$>>, which enables any $x \in S$ to decide which of $y, z \in S \setminus \{x\}$ is more similar to $x$. This “triplet comparison” has more general application than similarity induced by a symmetric distance function, as we discuss in [4], but disallows optimizations based on symmetric numerical similarity functions, used in the works cited above.

(b) **Stopping criterion:** Termination depends on a statistical criterion, presented in Section 3.3, applied to a sampled quantity called the friend clustering rate. By contrast, other implementations continue until no further updates are possible, except for a variant by [7] which stops when no more than a proportion $\delta$ of points allow updates.
(c) Parallelism: By casting the algorithm into a functional programming framework, we enable the Java Virtual Machine to distribute tasks among threads via a fork join pool, invisible to the programmer. Speedup due to parallelism is reported below.

2. K-nearest neighbors based on Comparators

2.1. Setting. Input data consist of a set $S$ of $n$ objects of type $V$, and what we call a ranking system [3], which attaches to each $x \in S$ a total order $\prec_x$ on $S \setminus \{x\}$; here $y \prec_x z$ is interpreted to mean that $y$ is more similar to $x$ than $z$ is. In data science, this is called a triplet comparison. The computer science equivalent is a Function<V, Comparator<V>>. Here we map each $x \in S$ to a specific Comparator [10], whose compare method depends on $x$. Formally compare($x; y, z$) < 0 means $y \prec_x z$, and compare($x; y, z$) > 0 means $z \prec_x y$, for distinct $x, y, z \in S$.

Such a family of Comparators gives an orientation of the line graph $L(K_n)$, called the ranking digraph. The relation $y \prec_x z$ is interpreted as an arc $xy \rightarrow xz$. See Figure 1 for an example. No comparison between $xy$ and $zw$ is provided if $\{x, y, z, w\}$ is a set of size four.

Most authors study the special case of a metric $\rho$ on $S$, where $y \prec_x z$ means $\rho(x, y) < \rho(x, z)$. Thus a metric (without ties) gives a total order (by distance) on points of the line graph $L(K_n)$.

Theorem: (Baron and Darling [3, Lemma 5.5]) The ranking digraph is acyclic if and only if the Comparators arise from some metric.

2.2. Example of non-metrizable Comparator. For points $x, y, z$ in the interior of the $d$-dimensional simplex (here $d \geq 3$), choose this Comparator: $y$ is closer to $x$ than $z$ is when

$$D(x\|y) < D(x\|z),$$

where $D(x\|y) := \sum_1^d x_i \log (x_i/y_i)$ is Kullback-Leibler (KL) divergence. KL divergence is asymmetric in the pair $(x, y)$, and does not satisfy the triangle inequality. This example was chosen to dispel the notion that a metric is needed. An example of a ranking digraph from six randomly generated points is shown in Figure 1. The existence of a $3$-cycle shows that these Comparators are not metrizable. Further examples and properties of ranking digraphs are described in [4].

2.3. K-NN graph. The K-NN graph is the directed graph with an arc from each $x \in S$ to the $K$ elements of $S \setminus \{x\}$ most similar to $x$. Naive computation would take $n$ calls to a Comparator-based sorting operation on $n-1$ objects, which would be $O(n^2 \log n)$ work – or in the special case of a metric, $\binom{n}{2}$ distance evaluations followed by an $O(n^2 \log n)$ sort. An ideal outcome would be to approximate the K-NN graph using $O(n \log K)$ calls to a Comparator-based sort of $O(K^2)$ objects.

Before we describe how K-NN Descents works, we mention that [11] lists half a dozen data science algorithms besides K-NN Descent that accept triplet comparisons as input, including another nearest neighbor search algorithm [2].

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1By definition, this relation is anti-symmetric and transitive for all $x$.

2An orientation of a graph $G$ is a digraph obtained by replacing each edge $\{a, b\}$ by one of the arcs $a \rightarrow b$ or $b \rightarrow a$. The textbook [2] explains graph-theoretic terms.

3Undirected edges of the complete graph $K_n$ on $S$ form the points of the line graph $L(K_n)$, in which $xy \sim xz$ for distinct $x, y, z$. Here $xy$ is an abbreviation for the edge $\{x, y\}$ of $K_n$. 
**Figure 1.** Ranking digraph on $\binom{6}{2}$ vertices, forming the unordered pairs from a set $\{x_i\}_{1 \leq i \leq 6}$, with each $x_i$ drawn uniformly at random from a 14-simplex. Vertex label $\{1,6\}$ (bottom right) refers to the unordered pair $\{x_1, x_6\}$. Its neighbors are $\{1,2\}, \{1,3\}, \{1,4\}, \{1,5\}$ together with $\{2,6\}, \{3,6\}, \{4,6\}, \{5,6\}$. Orientation of an arc such as $\{1,2\} \rightarrow \{1,4\}$ means that Kullback-Leibler divergences satisfy $D(x_1 \| x_2) < D(x_1 \| x_4)$. Proof by counterexample that this ranking system is not metrizable: vertices marked in black form a cycle $\{1,2\} \rightarrow \{1,4\}, \{1,4\} \rightarrow \{2,4\}, \{2,4\} \rightarrow \{1,2\}$.

3. **K-NN Descent Algorithm with a Statistical Stopping Rule**

3.1. **Friend-of-a-Friend Principle.** Dong, Charikar, and Li [7] base K-NN Descent on the Friend-of-a-Friend Principle, which states that a friend of a friend may be suitable as a friend. The algorithm consists of a sequence of rounds. At each round, we have a digraph on $S$, with regular out-degree $K$. Such a graph is called a $K$-out graph [8, Ch. 16]. If $x \rightarrow y$, call $y$ a **friend** of $x$, and $x$ a **co-friend** of $y$. If there is a ranking system on $S$, one $K$-out graph may be replaced by another $K$-out graph using a procedure we call **friend set update**.

Friend set update is like a cocktail party, attended by all members of $S$.

**Meet:** Each $x$ “meets” all friends of friends, and friends of co-friends, which become new acquaintances.

**Update:** The new friend set of $x$ consists of the closest $K$ out of all new acquaintances and former friends.
Table 2. Ingredients of the Java functional implementation of NND. For example, the Java object `rankingSystem` is an instance of the Java type `Function<V, Comparator<V>>`.

| Java Type                      | Instance Object               |
|--------------------------------|-------------------------------|
| `List<V>`                      | `points`                      |
| `Function<V, Comparator<V>>`   | `rankingSystem`               |
| `Map<V, Set<V>>`               | `friends, coFriends`          |
| `Function<V, Set<V>>`          | `proposeNewFriendSet`         |

3.2. Initialization of $K$-NN Descent.
- Each $x$ selects $K$ elements of $S \setminus \{x\}$ uniformly at random as its initial friend set. This digraph is called random $K$-out (Frieze & Karonski [8, Ch. 16]).
- Random $K$-out is an expander graph, whose undirected version has diameter $\leq \lceil \log_{K-1} n \rceil$ with high probability [3, Appendix].
- **Plausible heuristic:** $2\lceil \log_{K-1} n \rceil$ rounds of cocktail parties (friend updates) should suffice for “everyone to get to know each other”. Twice the diameter allows a “message” to travel from any vertex to its antipode, and back.

3.3. Sampling Method for Termination for $K$-NN Descent.
(i) Sample uniformly a point $x \in S$, and two friends $y, z \in S$ of $x$.
(ii) The friend clustering rate is the sample relative frequency that $y$ is a friend or cofriend of $z$.
(iii) The friend clustering rate is close to zero at the outset, and increases during the algorithm towards a plateau.
(iv) Stop at the first round at which the friend clustering rate does not increase, compared to the previous round.

4. A Java Implementation of $K$-NN Descent

4.1. Java parallel streams. The tools of modern Java [10], including generic types, pure functions, and parallel streams, enable a concise and performant distributed implementation of NND. Four main Java types are listed in the left column of Table 2 in monospaced font, and instances of these types used in NND are listed in the right column in boldface font.

Elements of $S$ have a generic type $V$, supplied by the invoking class (e.g. strings, vectors, trajectories). A ranking system is a `Function` from $V$ to a `Comparator` of objects of type $V$. Given three elements $x, y, z$ of $S$, the assertion that $y \prec x z$ is equivalent in Java to:

```
rankingSystem.apply(x).compare(y,z) < 0.
```

The `Comparator` for object $x$ is visible as an `x.getComparator()` method of the class $V$.

4.2. Distribution of tasks to processors. Initialize the `friends Map` so that the value associated with the key $x$ is a `Set<V>` object containing $K$ elements of $S \setminus \{x\}$ selected uniformly at random. The `coFriends Map` is derived from the `friends Map`.

Given $x$, the function `proposeNewFriendSet` (bottom of Table 2) compares all the cofriends, friends of friends, and friends of cofriends, with the current friend set of $x$, and proposes the best $K$ of all these as a new friend set. It is crucial that the friends of $x$ are not updated at the time the function is called. The `friends` and `coFriends` Maps remain
Table 3. Scaling observed in parallel streaming NND (dual Intel X5660, 12 cores total, JRE 11). Processing times decrease in successive rounds, because duplicate candidates are proposed by different friends. Time to execute a single round of NND scaled linearly with \(n\). When \(K\) was doubled, time to execute a round increased by a factor less than four. The FCC column shows the final value of the friend clustering coefficient, which appears to decrease with \(n\). The proportion of the true \(K\) nearest neighbors found, among a uniform sample of six points, was typically 95% or better; see Table 4. Results are consistent with the heuristic that \(2^{\frac{\log K}{n}}\) rounds of NND suffice.

| \(n\)     | \(K\) | rounds | \(2^{\log K / n}\) | 1st round | last round | FCC  |
|-----------|-------|--------|---------------------|-----------|------------|------|
| \(2 \times 10^4\) | 16    | 5      | 8                   | 1.9 sec   | 0.35 sec   | 0.271|
|           | 32    | 6      | 6                   | 2.6 sec   | 0.9 sec    | 0.264|
| \(2 \times 10^5\) | 16    | 7      | 10                  | 9.1 sec   | 5.3 sec    | 0.210|
|           | 32    | 5      | 8                   | 26 sec    | 13 sec     | 0.231|
| \(2 \times 10^6\) | 16    | 8      | 12                  | 88 sec    | 56 sec     | 0.205|
|           | 32    | 7      | 10                  | 295 sec   | 153 sec    | 0.210|
|           | 64    | 6      | 8                   | 1059 sec  | 401 sec    | 0.215|

effectively immutable while all these proposals are constructed in a parallel stream. This is part of the contract of java.util.stream, and makes it possible to execute a round of NND in a single line of code, by invoking the collect() method of Stream[10]:

```
points.parallelStream().collect(Collectors.toMap(x → x, x → proposeNewFriendSet.apply(x)));
```

The Map\(\langle V, \text{Set}\langle V\rangle\rangle\) produced by this command becomes the friends Map for the next round, and coFriends is updated accordingly.

The value type of the friends Map is NavigableSet\(\langle V\rangle\), with respect to the Comparator. Initial friends are ordered on insertion. During a friend set update at \(x\), each new candidate is compared to the last member of the NavigableSet at \(x\), and replaces it if appropriate.

The Java Virtual Machine allocates proposeNewFriendSet tasks among the processors and threads at run time. For example, experiments on a 12-core workstation with JRE 11 gave an eightfold speedup\(^4\) per round, compared to the same code where a single stream was used instead of a parallel stream. With \(n\) points and \(p\) processors, the speedup should be monotonically increasing in \(\frac{n}{p}\) for fixed \(K\), assuming that the limitation is thread contention for access to the friends and coFriends maps.

4.3. How many rounds of NND are needed? How many rounds of friend updates do we expect before the termination criterion of Section 3.3 is satisfied? Baron and Darling [3], and other citations therein, justify \(\lceil \log K / n \rceil\) as an estimate for the diameter of the undirected graph on \(S\) whose edges are the pairs \(\{x, y\}\), where \(y\) is an initial friend of \(x\). In our experiments, the number of rounds never exceeded, but was close to, \(2^{\lceil \log K / n \rceil}\); see Table 3.

4.4. Scaling of execution time with \(n\) and \(K\): The type \(V\) that we chose for our timing experiments was a point on the interior of the 9-dimensional standard simplex in \(\mathbb{R}^{10}\), representing a probability measure on a set of size ten. The ranking system was defined by taking

\(^4\)Here \(n = 2 \times 10^6, K = 16\). The speedup was much less for smaller \(n\).
Table 4. Here \( n = 2 \times 10^5 \) points were sampled from the \((d-1)\)-dimensional simplex in \( \mathbb{R}^d \), for four different values of \( d \), and \( K \)-NN descent was performed for \( K = 16 \) and \( K = 64 \). All runs finished within \( 2\lceil \log_K n \rceil \) (8 or 6) rounds. The FCC row shows the final value of the friend clustering coefficient. The accuracy row shows proportion of the true \( K \) nearest neighbors found, among a sample of six points. Note the gradual decline of accuracy with dimension, especially when the dimension \( d - 1 \) exceeds the number \( K \) of neighbors.

| \( K \) | feature | \( d = 10 \) | \( d = 20 \) | \( d = 40 \) | \( d = 60 \) |
|-------|---------|-------------|-------------|-------------|-------------|
| 64    | FCC     | 0.24        | 0.13        | 0.08        | 0.06        |
|      | accuracy| 1.0         | 1.0         | 0.90        | 0.84        |
| 16    | FCC     | 0.21        | 0.13        | 0.09        | 0.08        |
|      | accuracy| 0.95        | 0.52        | 0.43        | 0.36        |

\[ y \prec x z \] whenever 

\[ D(x\|y) < D(x\|z) \]

where \( D(x\|y) \) denotes Kullback-Leibler divergence of \( y \) from \( x \). The points themselves were sampled from a 10-dimensional Dirichlet distribution. Results are shown in Table 3 and discussed in the caption. The practical implications are:

1. A single call to \texttt{proposeNewFriendSet} costs \( O(K^2 \log K) \) work on average. Each round needs \( n \) calls to \texttt{proposeNewFriendSet}.
2. Run time for a single round of NND scales linearly with \( n \), for fixed \( K \).
3. The number of rounds of NND is not observed to exceed \( 2\lceil \log_K n \rceil \), suggesting an overall \( O((n \log n)K^2) \) run time, on cancelling two \( \log K \) factors.
4. For points on a 9-dimensional simplex, the accuracy (or recall) is 95% or better using our chosen stopping criterion, where accuracy means proportion of the true \( K \) nearest neighbors found, among a uniform sample of six points.
5. On \( p \) processors, parallel streams yield a speedup slightly less than \( p \), presumably because of thread contention. We observed at best an 8 times speedup on 12 cores.

4.5. Effect of dimension. The experiments in Table 3 were performed on points from a 9-dimensional simplex, taking \( K = 16, 32, 64 \). We also performed experiments where the points were drawn from \((d-1)\)-dimensional simplices, for \( d = 10, 20, 40, 60 \), comparing the cases \( K = 16 \) and \( K = 64 \). Table 4 shows a decline both in the friend clustering coefficient, and in the accuracy of the \( K \)-NN approximation as dimension increases, for fixed \( K \). Similar results are reported by Dong et al [7, Section 4.5], who interpret them as a consequence of the fact that, when sampling many points at random in high dimensions, the nearest neighbor and farthest neighbor of any point are at roughly the same distance [6].

5. Conclusions and future work

In the benign setting of probability measures sampled uniformly at random from a simplex in Euclidean space, with a comparator based on Kullback-Leibler divergence, performance of \( K \)-nearest neighbor descent conforms to the predictions based loosely on expander graphs.

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5On average \( K + 2K^2 \) items or fewer are proposed for insertion into a sorted set of size \( K \).

6We did not choose a larger sample size than six, because the random initialization already causes random variation in the outcome when NND is applied repeatedly to the same data set.
In particular, our statistical stopping criterion is satisfied within $2\lceil \log_K n \rceil$ rounds on a set of $n$ points, giving a run time proportional to
\begin{equation}
K^2 n \log n
\end{equation}
in contrast to the $O(n^{1.14})$ run time (for fixed $K$) reported by Dong et al [7]. The accuracy of NND in (intrinsic) dimension up to 20 is entirely satisfactory in examples studied, and does not require that the similarity measure be symmetric or derived from a metric.

Performance of NND can be much worse in other settings, such as a collection of long multi-character strings under a metric based on the longest common substring [3, Section 4.4]. More theory and new experiments will be needed to delineate the contexts in which $K$-nearest neighbor descent works well.

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