Efficient Parallel Algorithms for $k$-Center Clustering

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Abstract

The $k$-center problem is one of several classic NP-hard clustering questions. For contemporary massive data sets, RAM-based algorithms become impractical. And although there exist good sequential algorithms for $k$-center, they are not easily parallelizable.

In this paper, we design and implement parallel approximation algorithms for this problem. We observe that Gonzalez’s greedy algorithm can be efficiently parallelized in several MapReduce rounds; in practice, we find that two rounds are sufficient, leading to a 4-approximation. We contrast this with an existing parallel algorithm for $k$-center that runs in a constant number of rounds, and offers a 10-approximation. In depth runtime analysis reveals that this scheme is often slow, and that its sampling procedure only runs if $k$ is sufficiently small, relative to the input size. To trade off runtime for approximation guarantee, we parameterize this sampling algorithm, and find in our experiments that the algorithm is not only faster, but sometimes more effective. Yet the parallel version of Gonzalez is about 100 times faster than both its sequential version and the parallel sampling algorithm, barely compromising solution quality.

Keywords Clustering, $k$-center, approximation algorithms, parallel algorithms.

1 Introduction

Clustering is a fundamental task in interpreting data sets in contexts such as social networking, event recognition and bioinformatics. For many applications, the data sets can be prohibitively large, and there may be insufficient RAM to perform the necessary calculations efficiently, even when seeking approximate solutions. There are parallel-computing schemes such as MapReduce [7] that offer the ability to overcome the memory obstacle.

The $k$-center problem is a famous clustering problem, is NP-hard, and has well known (sequential) polynomial-time algorithms that offer essentially the best approximation possible. We describe a multi-round parallel algorithm for $k$-center, analyze in detail several parallel algorithms and compare them with one of these sequential methods. Inspired by theoretical guarantees and evaluation, we run comprehensive experiments, including trading off approximation for running time.
1.1 Clustering algorithms

Generally, clustering problems involve optimizing some function that indicates how well the clusters portray underlying structures in the data. In a metric clustering problem, the weights, representing the similarity between objects, observe the triangle inequality. The best-known example is of course the Euclidean metric. In the context of clustering, points in a metric space can be modelled as vertices in a complete graph. Each vertex stands for a data point, and each edge is weighted to indicate the distance (or dissimilarity) between the two adjacent points. The $k$-center problem is one of the fundamental NP-hard clustering problems on a metric input.

**Definition (k-CENTER).** *Find a set of at most $k$ centers — here we assume they are a subset of the vertices — such that the maximum distance from a vertex to its assigned center is minimized. The key task is to choose the optimum set of $k$ centers, as each of the remaining vertices would be assigned to its nearest center. For a set of points $V$, solution set $S$ containing at most $k$ vertices, and a distance function $d$, the objective of this problem can be considered as minimizing the objective $\min_{v \in V} \max_{s \in S} d(v, s)$.***

This objective has many applications, from vehicle routing to document clustering, in which it relates to concepts such as the furthest traveling time, or the least “similar” document. It can alternatively be considered to be minimizing the (maximum) covering radius of the clusters. Related classic NP-hard clustering questions include $k$-median, $k$-means, and facility location problems.

Via a reduction from the DOMINATING SET problem, Hsu and Nemhauser proved that, for all $\varepsilon > 0$, it is NP-hard to guarantee approximations within a factor $2 - \varepsilon$ of optimum for $k$-center [11]. Also exploiting the connection with the dominating set problem, Hochbaum and Shmoys gave a 2-approximation algorithm for the $k$-center problem [10]. Gonzalez introduced an greedy 2-approximation algorithm for the $k$-center problem [9]. Each of these $k$-center algorithms is inherently sequential, none admitting a simple parallel implementation.

1.2 Parallel algorithms

While approximation algorithms provide guaranteed performance with polynomial-time complexity, often data sets are large enough that running these algorithms efficiently requires prohibitively large amounts of RAM. For such instances, we can instead design algorithms that split the data across multiple machines, and process each part in parallel before aggregating the results. One important paradigm for parallel computing is MapReduce [7]. There are several fast approximation algorithms for famous clustering problems, such as $k$-center and $k$-median, in MapReduce [3, 6, 8]. Karloff et al. [13] introduced a theoretical model of computation for the MapReduce paradigm that is often applied to the analysis of MapReduce algorithms [2, 14]. They offered a comprehensive method for theoretically structuring algorithms for MapReduce, and defined a family of classes for MapReduce algorithms.

A MapReduce algorithm consists of a series of interleaving rounds of sequential *mappers* and parallel *reducers*. A map round assigns each data point independently to some reducer(s); the reducers run in parallel, each performing some procedure on the subset of points it has been assigned. A program in MapReduce may consist of several iterations of mappers and reducers, each involving potentially different map and reduce functions.

Both the $k$-means [2] and $k$-median [8] problems have been adapted to the MapReduce framework.

1.3 Our contribution

We provide a very careful and detailed examination both of the best-known MapReduce approximation algorithm for $k$-center [8], based on sampling, and a parallel implementation of Gonzalez’s algorithm (that
typically gives a 4-approximation). The 2-round special case of the latter approach was recently considered by Malkomes et al. [17], although their analysis and experiments differ considerably from ours. We describe in depth the performance and computational requirements of these approaches, and detail how this procedure can be adapted to allow for cases where RAM is insufficient even for the 2-round parallel solution. Based in part on a careful calculation of its running time, we generalize the sampling MapReduce scheme of Ene et al. [8], to trade off approximation guarantee for speed.

Our experiments show that the parallelized Gonzalez approach is often 100 times faster than the alternatives, while being almost as effective. These are the first experimental results for the k-center algorithm of Ene et al. [8]. Our results conform with the findings of Malkomes et al. [17], regarding the performance of their greedy approach.

2 Related work

2.1 Approximations

The k-center problem was first adapted to the MapReduce scheme by Ene et al. [8]. Their algorithm selects a uniformly random sample of the points, S, and adds points to S until most vertices of the graph are within a bounded distance of the sample. Finally, it adds remaining unrepresented vertices to S. A sequential k-center algorithm, with approximation factor $\alpha'$, is then run on S. With high probability, the k resulting centers constitute a $5\alpha'$-approximation for the k-center instance. When implemented using one of the 2-approximation algorithms described above, with high probability, this results in a 10-approximation overall. Ene et al. [8] apply a similar scheme to the k-median problem, with an $11\alpha'$-approximation (where $\alpha^*$ is a the factor of the standard approximation for k-median).

Recently, there has been increased interest in adapting k-center to MapReduce. Ceccarello et al. [5] gave a MapReduce diameter-approximation procedure with low parallel depth. From this, they derive a k-center solution for graphs with unit-weight edges: for $k \in \Omega(\log^2 n)$, with high probability, this is a $O(\log^3 n)$-approximation. Im and Moseley [12] have described a randomized 3-round 2-approximation algorithm that requires prior knowledge of the value of the optimal solution. Although they have announced that this leads to a 4-round 2-approximation without the requirement, the details have yet to be outlined. Very recently, Malkomes et al. [17] gave a 2-round approach similar to ours.

2.2 Experiments

Ene et al. [8] reported that their k-center MapReduce scheme performs poorly due to the sensitivity of k-center to sampling. Unfortunately, there are no results nor implementation details to confirm this. In combination with another simpler algorithm, we investigate the empirical performance their k-center scheme in greater detail.

Conversely, their k-median implementation performs significantly better than the worst-case guarantee. Solutions are comparable to sequential algorithms with much better bounds. Ene et al.’s results were based on the 3-approximation algorithm from Arya et al. [11]. There have been recent advances in k-median approximation algorithms, by Byrka et al. [4], as well as by Li and Svensson [15], and including these might improve the approximation bound.

3 Parallel k-center

We describe and analyze an approximation algorithm for the k-center problem that, for most practical cases, achieves a 4-approximation in only two MapReduce rounds. The intuition is that a sequential k-center algorithm finds in the first round a sample from each of the reducers such that the distance to all of the unsampled
points is bounded. Running a standard factor-2 algorithm on the sample reveals a factor-4 solution to the whole instance. Additional rounds can be performed in cases where even the sample is too large for a single machine: this would usually occur for very large values of $k$. Experiments show that this approach is often as good as that of the baseline sequential algorithm. The 2-round case of our algorithm is similar to the approach of Malkomes et al. [17]. Along with generalizing to larger instances, we analyze the run time of these algorithms in more detail, and provide an alternative, shorter proof of the two-round factor-four approximation.

3.1 Description

In this paper, the standard $k$-center approach is the factor-2-approximation of Gonzalez [9], which we refer to as GON. This algorithm chooses an arbitrary vertex from the graph, and marks it as a center. At each following step, the vertex farthest from the existing centers is marked as a new center, until $k$ centers have been chosen. As the edge weights comprise a metric, the triangle inequality ensures that the resulting set of centers comprises a 2-factor approximation.

Parallelized version Given a point set $V$ and a metric $d$, with $OPT$ representing the optimal covering radius, Algorithm 1 obtains a set of centers $\{c_i\}$ for which all points in $V_i$ – where $\{V_i\}$ partitions $V$ – are within radius $2 \cdot OPT$ from $c_i$. Running GON on $C = \bigcup_i c_i$ obtains $k$ centers whose covering radius for $C$ is $2 \cdot OPT$. Assume that we have $m$ machines each with capacity $c$. If $n/m \leq c$ and $k \cdot m \leq c$ then, due to the triangle inequality, this results in a 4-approximation MapReduce algorithm for $k$-center. If the sample is too large to fit onto the final machine, further iterations of the first round can be run on the sample until there are few enough points. Each additional round increases the approximation ratio by 2.

We dub this multi-round scheme for $k$-center MRG, for “MapReduce Gonzalez”, as shown in Algorithm 1.

Algorithm 1 MRG($V, k, m$)

1: $S \leftarrow V$
2: while $|S| > c$ do
3: The mapper arbitrarily partitions $V$ into sets $V_1, \ldots, V_m$ such that $\bigcup_i V_i = V$ and $|V_i| \leq \lceil n/m \rceil$, and each set $V_i$ is sent to a reducer $\rho_i$.
4: In parallel, each reducer $\rho_i$ runs GON on $V_i$, and returns a set $C_i$ containing the $k$ centers found.
5: $S \leftarrow \bigcup_i C_i$
6: The mapper sends all points in $S$ to a single reducer.
7: This reducer runs GON on $S$, and returns the set of centers $C^G$.
8: return $C^G$.

3.2 Approximation

Algorithm MRG clearly runs in polynomial time; to prove the four approximation of the 2-round case, we prove the following intermediate result. For an arbitrary subset $S$ of the vertex set $V$, let $S^G$ denote the set of points in the solution obtained by running GON on $S$, and let $SOL_S$ denote the value, the covering radius, of this solution.

Lemma 1. For each $S \subseteq V$, $SOL_S \leq 2 \cdot OPT$.

Proof. Let $V^*$ be an optimal set of centers. The vertex set $V$ can be partitioned into $k$ sets $\{V^*_j\}^m_{j=1}$ such that all points in set $V^*_j$ are within $OPT$ of some center $j \in V^*$.
First, assume every set $S^G \cap V^*_j$ has exactly one point. This point, $s_j$, can serve as the center for every point in $V^*_j$. Then every point $x$ in $V^*_j$, and hence in $V^*_j \cap S$, is within $2 \cdot OPT$ of $s_j$, as both $x$ and $s_j$ are within $OPT$ of $j \in V^*$.

However, if there is some partition $V^*_j$ with $|S^G \cap V^*_j| > 1$, then points in the same partition are within $2 \cdot OPT$ from each other. $GON$ adds a new center to $S^G$ only when it is the farthest from the points previously added to $S^G$. The presence of two centers within $2 \cdot OPT$ implies that all points in $S$ are within $2 \cdot OPT$ of $S^G$ (if there were some point farther, it would be in $S^G$ instead).

Therefore, for every subset $S$ of $V$, the value of the $k$-center solution returned by $GON$ on $S$ is at most twice the optimal solution for $V$. \hfill \Box

With sufficient space for $C$, the consequence of Lemma 1 is a factor-four approximation.

**Lemma 2.** If $n/m \leq c$ and $k \cdot m \leq c$, then the $k$-center algorithm can be implemented in MapReduce in two rounds with a 4-approximation guarantee.

**Proof.** Let $V_i$ refer to the points mapped to reducer $\rho_i$. Since we run $GON$ on $V_i$, every point in $V_i$ is within $2 \cdot OPT$ of a center in $C_i$ and hence in $C$.

According to Lemma 1, running $GON$ on $C$ arrives at a set of centers $C^G$ that is a $2 \cdot OPT$ solution on $C$. By the triangle inequality, it then follows that every vertex in the graph is within

\[ 2 \cdot OPT + 2 \cdot OPT = 4 \cdot OPT \]

of the $k$ centers $C^*$.

We now describe the properties of the setup and input for when $MRG$ can run effectively in two rounds. The capacity required is $O(\max(n/m, k \cdot m))$, based on which of the two rounds receives the most points. We assume that $n > k$, otherwise the solution to $k$-center is trivial. We further assume that $n/m > k$: if this is not the case, then we can reduce the number of machines. For small $k$, we only require that there is sufficient space across the machines to store the data set: that is, $m \cdot c \geq n$. We could also exploit external memory, for example by running multiple instances of our MapReduce algorithm and using a $k$-center algorithm on the disjoint union of the solutions; such cases are beyond the scope of this paper.

### 3.3 Multi-round analysis

If $k \cdot m > c$, we lack the required memory to store the sample on a single machine, and therefore run further iterations of the while loop. In such instances, $MRG$ uses more MapReduce rounds, loosening the approximation guarantee.

**Lemma 3.** If $n/m \leq c$ and $k \cdot m \geq c$, then the $k$-center problem can be implemented in $i$ rounds with a $2(i+1)$-approximation, where $i$ is chosen such that inequality 1 is satisfied.

During each round, the number of centers is decreased, ending when they fit on a single machine: each additional round adds two to the approximation factor. As $k \cdot m > c$ and $m \geq 1$, it follows that $k > c$. Even relaxing the requirement that $k \cdot m \leq c$, it is still necessary that $k \leq c$. Without this condition, selecting $k$ centers from a single machine seems to require incorporating external memory in some manner.

Assuming that $n/m \leq c$, after the first round we have $k \cdot m$ centers, so we send them to $m' = \lceil (k \cdot m)/c \rceil \leq (k \cdot m)/c + 1$ machines. After the second round, we have $k \cdot m'$ centers, which we can send to $m'' \leq \lceil (k \cdot m')/c \rceil \leq m \cdot k^2/c^2 + k/c + 1$ machines. In general, the number of machines required after $i$ rounds observes the bound

\[ m^{(i)} \leq m \cdot (k/c)^i + \frac{1 - (k/c)^i}{1 - k/c}, \]  

(1)
and we can run the final round when \( m^{(i)} < 2 \). As \( i \) increases, the second term in the inequality approaches \( 1/[1-(k/c)] \), which itself will be less than 2 only if \( 2k < c \). Intuitively, during each round we select \( k \) centers from each of the machines, so if \( k \) is close to \( c \) then the reduction in the number of centers in each round will be small.

4 Revisiting the sampling approach

In this section, we introduce a generalization of Ene et al.’s \([8]\) iterative-sampling procedure. As we show below, their algorithm is slower, but is more effective on the whole, than the sequential and parallel versions of Gonzalez’s algorithm. So that we can trade off runtime with approximation ratio, we add a new parameter to the iterative-sampling approach, and call this generalization EIM. Before this, we make some alterations to the scheme to prevent eccentric behaviors that sometimes occur.

4.1 Termination

The core of Ene et al.’s scheme is shown as Algorithm 2. Our implementation this algorithm adjusts the removal of points from \( R \) to ensure that the size of the set decreases in every iteration. For our implementation of line \([8]\) we remove vertices whose distance from \( S \) is equal to that from \( v \) to \( S \). In the original presentation such a vertex would remain in \( R \), which might lead to iterations in which no vertices are removed from \( R \), and the procedure looping indefinitely.

**Algorithm 2** EIM-MapReduce-Sample\((V,E,k,\epsilon)\)

1. \( S \leftarrow \emptyset, R \leftarrow V \)
2. while \(|R| > (4/\epsilon)kn^k \log n\) do
3. The mappers partition \( R \), and uniquely map each set \( R_i \) to a reducer \( i \).
4. Reducer \( i \) independently adds each point in \( R_i \) to set \( S' \) with probability \( 9kn^k(\log n)/|R| \), and to set \( H' \) with probability \( 4n^k(\log n)/|R| \).
5. Let \( H \leftarrow \cup_{1 \leq i \leq |R|} H' \) and \( S \leftarrow S \cup \left( \cup_{1 \leq i \leq |R|} S' \right) \). The mappers assign \( H \) and \( S \) to one machine, along with all edge distances between \( H \) and \( S \).
6. The reducer sets \( v \leftarrow \text{Select}(H,S) \).
7. The mappers arbitrarily partition \( R \), with \( R_i \) denoting these sets. Let \( v, R_i, S \), and the distances between \( R_i \) and \( S \) be assigned to reducer \( i \).
8. For \( x \in R_i \), remove \( x \) from \( R_i \) if \( d(x,S) \leq d(v,S) \).
9. Let \( R \leftarrow \cup_i R_i \).
10. Output \( C := S \cup R \).

With relatively small graphs, there is a non-trivial probability that the point \( v \in H \) will also be in \( S \). In such cases, the vertex \( v \) will be at equal distance to \( S \) as the points prior to it in the ordering given in line \([2]\) of \texttt{Select}() (Algorithm \([3]\)). This would mean that even points added to the sample might not be removed from \( R \). This increases the relative size of \( R \cap S \), also increasing the probability of no vertices being removed from \( R \) in subsequent rounds, as \( H \) is sampled from \( R \). If all points in \( R \) are eventually added to \( S \) then the algorithm cannot terminate. Therefore we assume that sampled points should always be removed from \( R \), and as such have adapted the algorithm to reflect this.

**Algorithm 3** Select\((H,S\)), with our parameter \( \phi \).

1. For each point \( x \in H \), find \( d(x,S) \)
2. Order the points in \( H \) according to their distance to \( S \) from farthest to smallest.
3. Let \( v \) be the point in position \( \phi(\log n)^{th} \) in the ordering.
4. return \( v \)
4.2 Trade-off

Ene et al. [8] prove that with high probability their MapReduce procedure runs in $O(1/\varepsilon)$ rounds. To decrease the number of rounds, we introduce parameter $\phi$ to $\text{Select()}$, which trades off approximation for running time. The original algorithm effectively fixed $\phi$ to be 8.

In the original EIM scheme, the expected number of points in $R$ that are farther from $S$ than $v$ is $8\log n \cdot |R|/|H| = |R| \cdot 2/n^\varepsilon$. By choosing a lower threshold for point $v$, we decrease the number of points that remain in $R$. Since the sampling algorithm terminates when $|R|$ falls below a the threshold defined by $v$, potentially this decreases the number of iterations. We introduce a variable $\phi$, and choose $v$ such that it is the $\phi(n)$th farthest point in $h$ from $S$.

To obtain a feasible $k$-center solution from the sample given by $\text{EIM-MapReduce-Sample()}$, a sequential $k$-center procedure is run on the resulting sample in an additional MapReduce round. Note that in line 3, $R$ is partitioned into $\lceil |R|/n^\varepsilon \rceil$ sets of size at most $\lceil n^\varepsilon \rceil$, and in line 7 the mappers partition $|R|/n^{1-\varepsilon}$ into $\lceil |R|/n^{1-\varepsilon} \rceil$ sets of size at most $\lceil |R|/n^{1-\varepsilon} \rceil$. In Section 6, we prove that – with weaker bounds, and with appropriate values of $\phi$ – the probabilistic 10-approximation still holds.

5 Runtime analysis

We now analyze in detail these parallel algorithms for $k$-center. Ene et al. [8] proved that their sampling procedure required $O(1/\varepsilon)$ rounds with high probability, while MRG can run in two rounds given sufficient resources. We consider also the computations required in each of the rounds to determine the expected overall runtime.

5.1 MRG

Assuming that $n/m \leq c$ and $k \cdot m \leq c$, MRG will run in two consecutive MapReduce iterations. The first iteration involves running $m$ concurrent $k$-center algorithms, each on $n/m$ vertices.

The runtime of $\text{GON}$ on $N$ points is $O(k \cdot N)$: each time a new center is selected, we need to find the distance of that center to all of the other vertices. So the runtime for the first round of MRG is $O(k \cdot n/m)$, with a low constant in the $O(\cdot)$ expression. In its second round, MRG runs $\text{GON}$ on the $k \cdot m$ centers obtained from the first round; this gives a runtime of $O(k^2 \cdot m)$. Therefore the total runtime of MRG is $O(k \cdot n/m + k^2 \cdot m)$, and for larger data sets, we would expect the dominant term to be $kn/m$.

5.2 EIM

The sampling algorithm, EIM, has, with high probability, $T \in \Theta(1/\varepsilon)$ iterations – each comprising three MapReduce rounds – followed by a final clean-up round at the end that solves a single $k$-center instance. Let $R_\ell$ and $S_\ell$ denote the state of sets $R$ and $S$, respectively, in iteration $\ell$ of the main loop of the algorithm. Counting from the first iteration, $|R_0| = n$ and, with high probability, $|R_\ell| = O(n/n^{\ell\varepsilon})$. In each iteration, points in $R$ are added to $H$ with probability $4n^\varepsilon (\log n)/|R|$, so $|H|$ is expected to be $O(n^\varepsilon \log n)$. And in line 5, $|S_\ell|$ becomes $|S_{\ell-1}| + O(kn^\varepsilon \log n)$, so that, starting with $|S_0| = 0$, we expect $|S_\ell| = O((\ell + 1)kn^\varepsilon \log n)$. We now analyze each MapReduce round.

**Round 1** (Lines 3 & 4). This round involves $O(|R_\ell|/m)$ operations during iteration $\ell$, so the total number of operations is

$$\sum_{\ell \in T} \frac{|R_\ell|}{m} \in O \left( \sum_{\ell \in T} \frac{n}{n^{\ell\varepsilon}} \right) \in O \left( \frac{1}{m} \cdot \frac{n}{1 - n^{-\varepsilon}} \right).$$
null
The sets at the core of EIM-MapReduce-Sample change with every iteration. Denote the state of sets \( R, S \) and \( H \) at the beginning of iteration \( \ell \) by \( R_{\ell}, S_{\ell}, \) and \( H_{\ell} \) respectively, where \( R_0 = V \) and \( S_0 = \emptyset \). The set of points that are removed from \( R \) during iteration \( \ell \) is denoted by \( D_\ell \), so \( R_{\ell+1} = R_\ell \setminus D_\ell \). Let \( U_\ell \) denote the set of points in \( R_\ell \) that are not satisfied by \( S_{\ell+1} \) with respect to \( Y \). Let \( U \) denote the set of all points that are not satisfied by \( SOL \) (the sample returned by the algorithm) with respect to \( Y \). If a point \( x \) is satisfied by \( S_\ell \) with respect to \( Y \), then it is also satisfied by \( SOL \) with respect to \( Y \), and therefore \( U \subseteq \bigcup_{\ell \geq 1} U_\ell \).

From the analysis of Ene et al. \[8\] we have the following lemma.

**Lemma 4.** Let \( Y \) be an arbitrary set with no more than \( k \) points. In iteration \( \ell \) of EIM-MapReduce-Sample, where \( \ell \geq 0 \),

\[
P[|U_\ell| \geq |R_\ell|/3n^\delta] \leq n^{-2}.
\]

We now show that our adaptation of the sampling algorithm by Ene et al. retains the same probabilistic 10-approximation guarantee.

**Lemma 5.** Let \( Y \) be a set of no more than \( k \) points. In iteration \( \ell \) of the while loop in EIM-MapReduce-Sample, let \( v_\ell \) denote the threshold in the current iteration: the point in \( H_\ell \) that is the \( \phi \log(n)^{th} \) most distant from \( S_{\ell+1} \). Then there exist values \( a \) and \( b \) such that, for some \( \gamma > 0 \),

\[
P\left[\frac{a|R_\ell|}{n^\delta} \leq |R_{\ell+1}| \leq \frac{b|R_\ell|}{n^\delta}\right] \geq 1 - \frac{2}{n^{1+\gamma}}.
\]

**Proof.** Recall that we selected a pivot point \( v \), and discarded the points that are well represented by the current sample, compared to \( v \). Note that \( R_{\ell+1} \) is the set of points in \( R_\ell \) such that the distance to the sample \( S_{\ell+1} \) is greater than the distance between the pivot point and the sample.

Ene et al. introduce these handy definitions. For a vertex \( t \), we refer to the number of points in \( R \) further from \( S_{t+1} \) than the point \( t \) as the \( \text{rank}_R \) of \( t \). For some value \( i \), and a set \( Y \subseteq R \), define \( L(i,Y) = |\{x \in Y : \text{rank}_R(x) \leq i\}| \) as the number of points in the set \( Y \) that have rank smaller than \( i \).

Let \( r = d|R_\ell|/n^\delta \), and let \( |H_\ell| = c \cdot n^\delta \log n \). By design,

\[
E[L(a \cdot r, H_\ell)] = a \cdot r \cdot |H_\ell|/|R_\ell| = acd \log n
\]

and

\[
E[L(b \cdot r, H_\ell)] = b \cdot r \cdot |H_\ell|/|R_\ell| = bcd \log n.
\]

If \( a \cdot r \leq |R_{\ell+1}| \leq b \cdot r \), then with high probability, the pivot (chosen to be the \( \phi \log n \) point in \( H_\ell \)) will be in the range \([L(a \cdot r, H_\ell), L(b \cdot r, H_\ell)]\).

By the Chernoff inequality,

\[
P[L(a \cdot r, H_\ell) \geq \phi \cdot \log n]
\]

\[
= P[L(a \cdot r, H_\ell) \geq (1 + \delta)E[L(a \cdot r, H_\ell)]]
\]

\[
= P[L(a \cdot r, H_\ell) \geq (1 + \delta) \cdot acd \log n]
\]

\[
\leq \exp\left[-\delta^2 \cdot acd \log n \over 2 + \delta\right]
\]

\[
= n^{-\delta^2 \cdot acd/(2 + \delta)}.
\]

Choosing \( \delta \) so that \((1 + \delta)E[L(a \cdot r, H_\ell)] \leq \phi \cdot \log n \) gives \( \delta \leq -1 + \phi/(acd) \). Since the Chernoff bound requires that \( \delta > 0 \), we insist that \( \phi > acd \).
The lemma statement requires $\Pr[L(a \cdot r, H_t) \geq \phi \cdot \log n] \leq n^{-(1+\gamma)}$, which we can achieve by finding values of $a$, $c$, $d$, and $\phi$ that for some $\gamma > 0$ satisfy
\[
\frac{(\phi/ (acd) - 1)^2 \cdot acd}{(2 + (\phi/ (acd) - 1))} \geq (1 + \gamma).
\]
Letting $x = 1 + \gamma$, this is equivalent to $(acd)^2 - (2\phi + x)acd + (\phi^2 - x\phi) \geq 0$, which has real roots at $acd = \phi + x/2 \pm \sqrt{2\phi x + x^2/4}$. Similarly,
\[
\Pr[L(b \cdot r, H_t) \leq \phi \cdot \log n]
= \Pr[L(b \cdot r, H_t) \leq (1 - \delta)E[L(b \cdot r, H_t)]]
= \Pr[L(b \cdot r, H_t) \leq (1 - \delta) \cdot bcd \log n]
\leq \exp\left[\frac{-\delta^2 \cdot bcd \log n}{2}\right]
= n^{-\delta^2 bcd/2} \leq n^{-x}.
\]
Choosing $\delta$ so that $(1 - \delta)E[L(b \cdot r, H_t)] \leq \phi \cdot \log n$ gives $\delta \leq 1 - \phi / (bcd)$. Since the Chernoff bound requires that $\delta > 0$, this gives the constraint $\phi < bcd$.

For the last inequality to hold, we need to find values of $b$, $c$, $d$, and $\phi$ such that
\[(bcd)^2 - (2\phi + x)bcd + \phi^2 \geq 0.
\]
This has real roots at $bcd = \phi + x/2 \pm \sqrt{2\phi x + x^2/4}$.

This gives feasible solutions for $acd \leq \phi + x/2 - \sqrt{2\phi x + x^2/4}$ and $bcd \geq \phi + x + \sqrt{2\phi x + x^2}$. For later results we require that $a = 1$ and $b \leq 5$. So for there to exist feasible values of $c$ and $d$, we have the following constraint,
\[
\frac{\phi + x + \sqrt{2\phi x + x^2}}{b} \leq \phi + \frac{x}{2} - \sqrt{2\phi x + \frac{x^2}{4}},
\]
where $b \leq 5$ and $x = 1 + \gamma$. When this bound holds, we can find values of each of the parameters such that the probability of $|r_{\ell+1}|$ being outside of the defined bounds is less than $1/n^\gamma$, and therefore
\[
\Pr\left[\frac{|R_{\ell}|}{n^{\varepsilon}} \leq |R_{\ell+1}| \leq \frac{5|R_{\ell}|}{n^{\varepsilon}}\right] \geq 1 - 1/n^{\gamma} - 1/n^\gamma,
\]
which is $1 - 2n^{-(1+\gamma)}$. With this probability, the number of points in $R_{\ell}$ that are further from $S$ than $v_{\ell}$ (and hence the size of the set $R_{\ell+1}$) is in the range $[|R_{\ell}|/n^{\varepsilon}, 5|R_{\ell}|/n^{\varepsilon}]$. \hfill \qed

Ene et al. \cite{ene2015approximate} prove that with probability $1 - O(1/n)$, it is possible to map each unsatisfied point to a satisfied point such that no two unsatisfied points are mapped to the same satisfied point. Such a mapping allows them to bound the cost of the unsatisfied points with regards to the cost of the optimal solution. Their proof relies on the choice of $b = 4$, and the bound from Lemma \ref{lem:cd} giving a probability greater than $1 - 2n^{-2}$. However, we use $b \leq 5$, and only assure a probability of $1 - 2n^{-(1+\gamma)}$. Therefore, we prove that the required mapping exists with probability $1 - O(n^{-(1+\gamma)})$; by setting $\gamma = (\log \log n) / \log n$ this gives a probability of $1 - O(1/\log n)$, which is sufficient for large values of $n$. The choice of $b$ arises from the requirement that $b/n^{\varepsilon} < 2$: for $\varepsilon = 0.1$ and $n \geq 10,000$, this holds for $b \leq 5$.

In the original analysis, Ene et al. proved that their results hold with high probability, which they define as having probability $\geq 1 - O(1/n^{2\gamma})$. We instead bound our confidence in these results with probability $1 - O(1/n^{1+\gamma})$ for a variable $\gamma$, which we will refer to as with sufficient probability, or w.s.p.

The following results follow from the above analysis and that given by Ene et al. \cite{ene2015approximate}.
Table 1: Theoretical comparison of algorithms: Approximation factor represented by $\alpha$, run times are asymptotic, $O(\cdots)$.

| Algorithm | $\alpha$ | Rounds | Runtime                                |
|-----------|----------|--------|----------------------------------------|
| GON [9]   | 2        | n/a    | $k \cdot n$                            |
| MRG       | 4        | 2      | $kn/m + k^2m$                          |
| EIM [8]   | 10       | $O(1/\epsilon)$ | $kn^{1+\epsilon} \log n \over m(1-n^{-\epsilon})^2$ |

Lemma 6. For the sample $S$ returned by EIM-MapReduce-Sample, w.s.p. we have $OPT(V, S) \leq 5 \cdot OPT$.

Lemma 7. The procedure resulting from running an $\alpha$-approximation algorithm on the sample returned by EIM-MapReduce-Sample achieves a $4\alpha + 2$-approximation for the $k$-centre problem w.s.p.

When running a 2-approximation algorithm on the sample, this result gives a 10-approximation bound on the resulting procedure. To achieve a success probability higher than $1 - O(1/n)$, we need $x \geq 1$: by the bound in Inequality (2), this implies that $\phi > 5.15$.

7 Experiments

In this section we compare three algorithm (families), both in terms of speed and effectiveness, and contrast the theoretical properties of these methods (as shown in Table 1) with their empirical performance.

7.1 Setup

We run experiments on three algorithm families, each of which we implement in the C language. First is the typically 2-round algorithm, MRG; second is (our Section 6 generalization of) the sampling algorithm of Ene et al. [8], EIM; third is the (standard) sequential algorithm, GON. The latter, with its factor-2 approximation guarantee serves as an effectiveness baseline.

For the sake of consistency with previous literature, our method of implementing these algorithms mimics that of Ene et al. [8] in several ways. In particular, we adopt a MapReduce approach, but do not record the cost of moving data between machines. (As MRG involves fewer rounds, the expected cost of moving data between machines would be less than for EIM.) We simulate the parallel machines sequentially on a single machine, taking the longest processing time of the simulated machines as the processing time for that MapReduce round. For all parallel implementations, GON is the subprocedure for selecting the final centers.

The experimental system is a ‘commodity’ machine, with 8GB of main memory and an Intel® Core™ i7-2600 CPU @ 3.40GHz.

7.2 Experimental design

Ene et al. [8] generated synthetic data, designed to have a fixed number of similarly sized clusters. Moreover, they tested their algorithm for values of $k$ equal to the number of clusters. We evaluate the algorithms over a range of values of $k$ and vary the numbers of inherent clusters. In practice, the number of clusters may not be known in advance, and the number of clusters required can be independent of the structure of the data. To better determine how well these algorithms are likely to perform in practice, we extend these experiments to test on graphs with different underlying structures.

In all of the experiments, the distance is Euclidean, computed as required from the locations of the points. The $k$-center algorithm assumes a complete graph as input, and a matrix representation of a graph,
with all distances stored explicitly, might result in a significant proportion of the data sent between machines being unnecessary. The number of machines, \( m \), is fixed to 50, while \( n \) and \( k \) vary. Our preliminary experimentation with the EIM algorithm, over a range of values of \( \epsilon \), confirms that Ene et al.'s choice of \( \epsilon = 0.1 \) was good.

In Section 6 we introduced a parameter \( \phi \) to the EIM sampling approach. In our experiments, we test the effect of lowering \( \phi \) from its “original” value of 8, both in terms of runtime and effectiveness. Corresponding with our theoretical results in Section 6 we choose \( \phi = 6 \); and to determine the robustness of the algorithm, we test with \( \phi = 4 \) and \( \phi = 1 \), which are below the bound of \( \phi = 5.15 \) that was given in Section 6.

### 7.3 Data sets

We test against a combination of real and synthetic data sets, primarily in two and three dimensions, but with several real data sets of larger dimension. The data sets have a range of sizes, from 10,000 through to 1,000,000 points, with varying degrees of inherent clustering. Our synthetic data sets have three different formats, viz.

**UNIF** The \( n \) points are uniformly distributed in a two-dimensional square.

**GAU** The \( k' \) cluster centers – where \( k' \) might not equal \( k \) – are uniformly randomly generated in a unit cube. The \( n \) points are distributed into these clusters uniformly at random, resulting in clusters of roughly similar size. This helps determine the accuracy with which the procedures can identify different clusters. Distance from points to the cluster center follows a Gaussian distribution with \( \sigma = 1/10 \). These data sets mimic those used in the experiments of Ene et al [8].

**UnB** An unbalanced arrangement, similar to GAU. The distribution of points to inherent clusters is biased such that around half of the points are in a single (inherent) cluster; the distribution between the remaining clusters remains uniform.

We generate three graphs of each size and type, and run the algorithms twice over each data set, taking the average. This gives a total of six results for each type of data set, over three different graphs.

We take real data sets from the UCI Machine Learning Repository [16], over a wide range of sizes, applications and dimensions. We run four tests over each of the real data sets, and take the average result. We include results for the 25,010-point training set for the POKER HAND data set, and the 10% sample from the 4,000,000-point KDD CUP 1999 data set.
8 Results

Overall MRG is faster than the alternative procedures - often by orders of magnitude, with EIM running slower than the sequential algorithm despite being parallelized, conforming with the analysis in Section 5.

8.1 Summary

In most cases, despite having worse approximation guarantees, the solutions for the parallelized algorithms are comparable to those of the baseline, GON, with EIM performing slightly better for synthetic data sets. Ene et al [8] suggested that their sampling-based algorithm did not perform particularly well, likely due to the k-center procedure being sensitive to outliers. Our experimental results show otherwise: sampling fewer points can occasionally provide better results due to the tendency to avoid sampling points that are well represented, but toward the edge of the cluster. The tendency for GON to favor outliers is often mitigated, rather than amplified, by sampling. As shown in Table 4, this effect is particularly evident for GAU graphs where $k = k'$.
As illustrated in Tables 2 and 4 for the synthetic data sets, the parallel algorithms are about as effective as Gonzalez’s algorithm. In general, EIM is slightly more effective than MRG. With the exception of the EIM results on the KDD CUP 1999 10% sample, for which it performs poorly, the same occurs on the real data sets, as seen in Figure 1 and Table 5.

### Table 4: Solution value over k for UnB (n = 200,000, k’ = 25). When k = k’, EIM is notably better.

| k  | MRG  | EIM  | GON |
|----|------|------|-----|
| 2  | 97.96| 93.69| 93.37|
| 5  | 64.61| 64.28| 61.72|
| 10 | 40.17| 40.05| 40.39|
| 25 | 0.932| 0.828| 0.939|
| 50 | 0.668| 0.643| 0.655|
| 100| 0.515| 0.530| 0.500|

### Table 5: Solution value over k for the Poker Hand data set.

| k  | MRG  | EIM  | GON |
|----|------|------|-----|
| 2  | 19.41| 18.60| 18.17|
| 5  | 18.06| 17.07| 17.25|
| 10 | 15.12| 14.20| 15.03|
| 25 | 12.13| 11.98| 11.84|
| 50 | 10.07| 9.418| 9.617|
| 100| 8.774| 9.241| 8.396|

As illustrated in Tables 2 and 4, for the synthetic data sets, the parallel algorithms are about as effective as Gonzalez’s algorithm. In general, EIM is slightly more effective than MRG. With the exception of the EIM results on the KDD CUP 1999 10% sample, for which it performs poorly, the same occurs on the real data sets, as seen in Figure 1 and Table 5.

### 8.2 Running time

For the majority of the experiments, EIM ran using two iterations of the main loop, for a total of seven MapReduce rounds. On certain data sets, EIM sometimes executes one iteration, sometimes two – that is, four or seven MapReduce rounds – as the number of points removed per round is probabilistic.

From Figures 2b, 3b, and 4b we can see that as the ratio of n to k drops, at some point, EIM merely sends the entire data set to a single machine, rather than employing the sampling procedure. We can also...
Table 6: Average solution value over $\phi$, in EIM, for GAU ($n = 200,000, k' = 25$). For each $k$, the lowest value is in *italics*.

| $k$ | $\phi$ | 1  | 4  | 6  | 8  |
|-----|--------|----|----|----|----|
| 2   | 88.4   | 80.4| 85.5| 86.5|    |
| 5   | 59.9   | 60.9| 56.5| 61.9|    |
| 10  | 36.2   | 35.5| 34.7| 35.3|    |
| 25  | 0.796  | 0.780| 0.826| 0.840|    |
| 50  | 0.630  | 0.617| 0.610| 0.666|    |
| 100 | 0.478  | 0.492| 0.505| 0.535|    |

Table 7: Average runtime over $\phi$, in EIM, for GAU ($n = 200,000, k' = 25$). The lowest runtime in each row is in *italics*.

| $k$ | $\phi$ | 1  | 4  | 6  | 8  |
|-----|--------|----|----|----|----|
| 2   | 0.050  | 0.059| 0.165| 0.135|    |
| 5   | 0.080  | 0.130| 0.368| 0.314|    |
| 10  | 0.283  | 0.480| 0.549| 0.552|    |
| 25  | 0.588  | 0.505| 1.47 | 1.42 |    |
| 50  | 0.693  | 0.816| 2.84 | 2.24 |    |
| 100 | 0.726  | 0.757| 3.78 | 3.59 |    |

note that in Figure 4b, MRG displays a different trend from Figure 4a. In Section 5 we showed that the runtime is $O(kn/m + k^2 \cdot m)$. For larger values of $k$ and small values of $n$, the $k^2 \cdot m$ term dominates; as $n$ grows, the $k \cdot n/m$ term dominates, so the trend becomes similar to that in Figure 4a. From our analysis in Section 5 both MRG and EIM have a round with a $k^2$ term in the running time. When $k$ is large relative to $n$, this can potentially dominate.

8.3 Runtime/Approximation Trade-off

We examine the sensitivity of the EIM algorithm to the $\phi$ parameter. As expected, the variability of effectiveness increases as the $\phi$ parameter decreases, while the runtimes significantly decrease. Tables 6 and 7 compare the average solution value and runtimes for the different parameters. The algorithm speeds up significantly for values of $\phi$ below the threshold of 5.15 (above which there is a guaranteed low probability of poor solutions, see Section 6). Yet, in practice, it still returns acceptable solutions: in some cases solutions are even better with smaller values of $\phi$.

This seemingly counterintuitive behavior can be explained by the choice of GON as the sub-procedure for the sample. As noted above, selecting the farthest points as new centers makes it more likely that points at the perimeter of a cluster are chosen; although each cluster is well represented by some vertex, the selected center is at the perimeter of the cluster. By sampling fewer points, it is less likely that points that are extremal to the cluster are present in the subgraph on which GON is run. Therefore in decreasing the runtime of the algorithm, for appropriate values of $k$, we can potentially improve the average value of the solutions obtained. However this behavior is likely to be more volatile: the guaranteed bound on the performance has lower probability, giving a higher chance that a very poor solution is returned.
Figure 4: Runtimes in seconds for fixed $k$ over values of $n$ ranging from $10,000$ to $1,000,000$. For sufficiently small values of $n$ relative to $k$, EIM behaves identically to GON. The is caused by the condition on the while loop: if $k$ is large enough, the condition is never met and no sampling occurs, so GON is run on the entire data set.

9 Conclusion

In this paper, we describe a multi-round parallel procedure for the $k$-center problem. When it runs in only 2 MapReduce rounds, it is a 4-approximation. We show experimentally that it returns solutions that compare well to those of a sequential 2-approximation algorithm, while running extremely fast.

We compare this approach to the existing 10-approximation sampling-based MapReduce procedure [8]. It is often slightly more effective, but can be very slow. To support our experimental results, we give the first detailed runtime analysis for the sampling approach, the proof of which correspond with our empirical results. We also parameterized the sampling procedure to improve runtimes, sometimes even bringing better solutions despite the lack of a provable effectiveness bound.

Future work

The approximation factor of four for MRG is tight. There are graphs on which, with adversarial assignment of points to machines and choice of seedings for GON, MRG gives a 4-approximation. How likely such cases are in practice? We seek bounds on the probability that this algorithm gives a poor approximation. And what is the effectiveness when MRG needs more than two rounds?

Recently, Im and Moseley [12] described a 3-round 2-approximation MapReduce procedure for the $k$-center problem under the assumption that $OPT$ is known, and announced a 4-round procedure that does not require prior knowledge of the optimal solution – these details have yet to appear. More recently Malkomes et al. [17] presented a parallel adaptation of the $k$-center algorithm comparable to a special case of our approach. Currently all such approaches rely on the sequential algorithm of Gonzalez [9]. It would be interesting to compare with similar adaptations of alternative sequential algorithms, such as that of Hochbaum & Shmoys [10].

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