# **Bandit-PAM:** Almost Linear Time k-Medoids Clustering via Multi-Armed Bandits

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#### **Abstract**

Clustering is a ubiquitous task in data science. Compared to the commonly used k-means clustering algorithm, k-medoids clustering algorithms require the cluster centers to be actual data points and support arbitrary distance metrics, allowing for greater interpretability and the clustering of structured objects. Current state-of-theart k-medoids clustering algorithms, such as Partitioning Around Medoids (PAM), are iterative and are quadratic in the dataset size, n, for each iteration, prohibitively expensive for large datasets. We propose Bandit-PAM, a randomized algorithm inspired by techniques from multi-armed bandits, that significantly improves the computational efficiency of PAM. We theoretically prove that Bandit-PAM reduces the complexity of each PAM iteration from  $O(n^2)$  to  $O(n \log n)$  and returns the same results with high probability, under assumptions on the data that often hold in practice. We empirically validate our results on several large-scale real-world datasets, including the a coding exercise submissions dataset from Code.org, the 10x Genomics 68k PBMC single-cell RNA sequencing dataset, and the MNIST handwritten digits dataset. We observe that Bandit-PAM provides returns the same results as PAM while performing up to 200x fewer distance computations. The improvements demonstrated by Bandit-PAM enable k-medoids clustering on a wide range of applications, including identifying cell types in large-scale single-cell data or providing scalable feedback for students learning computer science.

### 1 Introduction

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21 Many modern data science applications require the clustering of very-large-scale data. Due to its 22 computational efficiency, the *k*-means clustering algorithm [27, 25] has been one of the most widely-23 used clustering algorithms. *k*-means alternates between assigning points to their nearest cluster 24 centers and recomputing those centers. Central to its success is the specific choice of the cluster 25 center: for a set of points, *k*-means defines the cluster center as the point with the smallest average 26 squared Euclidean distance to all other points in the set. Under such a definition, the cluster center is 27 the arithmetic mean of the cluster's points and can be computed efficiently.

While commonly used in practice, k-means clustering suffers from several drawbacks. First, while one can efficiently compute the cluster centers under squared Euclidean distance, it is not straightforward to generalize to other distance metrics [32, 12, 5]. However, a different distance may be desirable in different applications. For example,  $l_1$  and cosine distance are often used in sparse data, such as in recommendation systems [23] and single-cell RNA-seq analysis [31]; additional examples include string edit distance in text data [29], and graph metrics in social network data [28]. Second, the cluster center in k-means clustering is in general not a point in the dataset and may not be interpretable in many applications. This is especially problematic when the data is structured, such as parse trees in

context-free grammars, sparse data in recommendation systems [23], or images in computer vision where the mean image is visually random noise [23].

Alternatively, *k*-medoids clustering algorithms [16, 17] use the *medoid* to define the cluster center for a set of point, where for an arbitrary distance function, the medoid is the point *in the set* that minimizes the average distance to all the other points. Note that the distance metric can be arbitrary—indeed, it need not be a distance metric at all and could be an asymmetric dissimilarity measure—which addresses the first shortcoming of *k*-means outlined above. Moreover, unlike *k*-means, the cluster centers in *k*-medoids, i.e. the medoids, are restricted to be points in the dataset, thus addressing the second shortcoming of *k*-means clustering described above.

Despite its advantages, k-medoids clustering is less popular than k-means due to its computational efficiency. Indeed, state-of-art k-medoids clustering algorithms are iterative and are quadratic in the data size, whereas k-means is linear in dataset size in each iteration.

Mathematically, for n data points  $\mathcal{X}=\{x_1,x_2,\cdots,x_n\}$  and a user-specified distance function  $d(\cdot,\cdot)$ , the k-medoids problem is to find a set of k medoids  $\mathcal{M}=\{m_1,\cdots,m_k\}\subset\mathcal{X}$  to minimize the overall distance of points from their closest medoids:

$$L(\mathcal{M}) = \sum_{i=1}^{n} \min_{m \in \mathcal{M}} d(m, x_i)$$
(1)

This problem is, unfortunately, NP-hard in general [35]. Partitioning Around Medoids (PAM) [16, 17] is one of the most widely used heuristic algorithms for k-medoids clustering. PAM is split into two subroutines: BUILD and SWAP. First, in the BUILD step, PAM aims to find an initial set of k medoids by greedily and iteratively selecting points that minimize the k-medoids clustering loss as described in (1). Next, in the SWAP step, PAM considers all k(n-k) possible pairs of medoid and non-medoid points and swaps the pair that reduces the loss the most. The SWAP step is repeated until no further improvements can be made.

PAM has been empirically shown to produce better results than other popular k-medoids clustering algorithms [34, 35]. However, the BUILD step and each of the SWAP steps require  $O(kn^2)$  distance evaluations and can be prohibitively expensive to run, especially for large datasets or when the distance evaluations are themselves expensive (e.g. edit distance between two long strings).

Randomized algorithms like CLARA [17] and CLARANS [30] have been proposed to improve computational efficiency, but at the cost of deteriorated clustering quality. More recently, Schubert et al. [35] proposed a deterministic algorithm, dubbed FastPAM1, that guarantees the same output as PAM but improves the complexity to  $O(n^2)$  when the cluster sizes are similar. However, the factor O(k) improvement becomes less relevant when the sample size n is large and the number of medoids k is relatively small compared to n. Throughout the rest of this work, we treat k fixed.

Contributions: In this work, we propose a novel randomized k-medoids algorithm, called Bandit-PAM, that significantly improves the computational efficiency of PAM while returning the same result with high probability. We theoretically prove that Bandit-PAM reduces the complexity on the sample size n from  $O(n^2)$  to  $O(n \log n)$ , both for the BUILD step and each SWAP step, under reasonable assumptions that hold in many real-world datasets. We empirically validate our results on several large-scale real-world datasets and observe that Bandit-PAM provides up to 200x reduction of distance computations while returning the same result as PAM. We also release a high-performance C++ of implementation Bandit-PAM, which brings a 3.2x wall-clock-time speedup over the state-of-the-art FastPAM implementation [36] on the full MNIST dataset without precomputing and caching the the  $n^2$  pairwise distance matrix as in previous approaches.

Intuitively, Bandit-PAM works by recasting each step of PAM from a *deterministic computational* problem to a *statistical estimation problem*. In the BUILD step assignment of the lth medoid, for example, we need to choose the point amongst all n-l non-medoids that will lead to the lowest overall loss (1) if chosen as the next medoid. Mathematically, we wish to find x that minimizes:

$$L(x; \mathcal{M}) = \sum_{j=1}^{n} \min_{m \in \mathcal{M} \cup \{x\}} d(m, x_j) \stackrel{\text{def}}{=} \sum_{j=1}^{n} g(x_j), \tag{2}$$

where  $g(\cdot)$  is a function that depends on  $\mathcal{M}$  and x. Eq. (2) shows that the loss of a new medoid assignment  $L(\mathcal{M};x)$  can be written as the summation of the value of the function  $g(\cdot)$  evaluated

on all n points in the dataset. Though approaches such as PAM compute  $L(\mathcal{M};x)$  exactly for each x, Bandit-PAM adaptively estimates this quantity by sampling reference points  $x_i$  for the most promising candidates. Indeed, computing  $L(\mathcal{M}; x)$  exactly for every x is not required; promising candidates can be estimated with higher accuracy (more reference point  $x_i$ 's) and less promising ones can be discarded early without requiring further unnecessary computation.

To design the adaptive sampling strategy, we show that the BUILD step and each SWAP iteration can 89 be formulated as a best-arm identification problem from the multi-armed bandits (MAB) literature 90 [1, 10, 13, 14]. In the typical version of the best-arm identification problem, we have m arms. At each 91 time step  $t=0,1,\cdots$ , we decide to pull an arm  $A_t\in\{1,\cdots,m\}$ , and receive a reward  $R_t$  with 92  $E[R_t] = \mu_{A_t}$ . The goal is to identify the arm with the largest expected reward with high probability 93 while expending the fewest number of total arm pulls. In the BUILD step, we view each candidate 94 medoid x as an arm in a best-arm identification problem. The arm parameter corresponds E[g], and 95 by pulling an arm, we observe the loss evaluated on a randomly sampled data point  $x_i$ . Using this 96 reduction, the best candidate medoid can be estimated using existing best-arm algorithms like the Upper Confidence Bound (UCB) algorithm [21].

**Related work:** Many other k-medoids algorithms exist, in addition to CLARA, CLARANS, and FastPAM as described above. Park et al. [33] proposed a k-means-like algorithm that alternates between reassigning the points to their closest medoid and recomputing the medoid for each cluster until the k-medoids clustering loss can no longer be improved. Other proposals include optimizations for Euclidean space and tabu search heuristics [9]. Recent work has also focused on distributed PAM, where the dataset cannot fit on one machine [37]. All of these algorithms, however, scale quadratically in dataset size and could benefit from improvements in complexity.

The idea of algorithm acceleration by converting a computational problem into a statistical estimation 106 problem and designing the adaptive sampling procedure via multi-armed bandits has witnessed a 107 few successes [7, 18, 24, 15, 3, 39]. In the context of k-medoids clustering, previous work [2, 4] has 108 considered finding the single medoid of a set points (i.e. the 1-medoid problem). In these works, the 109 1-medoid problem was also formulated as a best-arm identification problem, with each point being an arm and its average distance to other points being the arm parameter. 111

While the 1-medoid problem considered in prior work can be solved exactly, the k-medoids problem 112 is NP-Hard and is therefore only tractable with heuristic solutions. Hence, this paper focuses on improving the computational efficiency of an existing heuristic solution, PAM, that has been empirically observed to be superior to other techniques. Moreover, instead of having a single best-arm identification problem, we reformulate PAM as a sequence of best-arm problems. We treat different objects as arms in different steps of PAM; in the BUILD step, each point corresponds to an arm, whereas in the SWAP step, each medoid-and-non-medoid pair corresponds to an arm. We further 119 notice that the intrinsic difficulties of this sequence of best-arm problems are different, which can be exploited to further speed up the algorithm, as demonstrated in Section 5.

#### **Preliminaries**

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For n data points  $\mathcal{X} = \{x_1, x_2, \dots, x_n\}$  and a user-specified distance function  $d(\cdot, \cdot)$ , the k-medoids 122 problem aims to find a set of k medoids  $\mathcal{M} = \{m_1, \dots, m_k\} \subset \mathcal{X}$  to minimize the overall distance 123 of points from their closest medoids: 124

$$L(\mathcal{M}) = \sum_{i=1}^{n} \min_{m \in \mathcal{M}} d(m, x_i)$$
(3)

Note that d does not need to satisfy symmetry, triangle inequality, or positivity. For the rest of the 125 paper, we use [n] to denote the set  $\{1, \dots, n\}$  and  $|\mathcal{S}|$  to represent the cardinality of a set  $\mathcal{S}$ . For two 126 scalars a, b, we let  $a \wedge b = \min(a, b)$  and  $a \vee b = \max(a, b)$ . 127

#### 2.1 Partitioning Around Medoids (PAM)

The original PAM algorithm [16, 17] first initializes the set of k medoids via the BUILD step and 129 then repeatedly performs the SWAP step to improve the loss (3) until convergence. 130

**BUILD:** PAM initializes a set of k medoids by greedily assigning medoids one-by-one so as to minimize the overall loss (3). The first point added in this manner is the medoid of all n points. Given the current set of l medoids  $\mathcal{M}_l = \{m_1, \dots, m_l\}$ , the next point to add  $m^*$  can be written as

BUILD: 
$$m^* = \underset{x \in \mathcal{X} \setminus \mathcal{M}_l}{\arg \min} \frac{1}{n} \sum_{j=1}^n \left[ d(x, x_j) \wedge \underset{m' \in \mathcal{M}_l}{\min} d(m', x_j) \right].$$
 (4)

SWAP: PAM then swaps the medoid-nonmedoid pair that would reduce the loss (3) the most among all possible k(n-k) such pairs. Let  $\mathcal M$  be the current set of k medoids. Then the best pair to swap is

SWAP: 
$$(m^*, x^*) = \underset{(m,x) \in \mathcal{M} \times (\mathcal{X} \setminus \mathcal{M})}{\arg \min} \frac{1}{n} \sum_{j=1}^{n} \left[ d(x, x_j) \wedge \underset{m' \in \mathcal{M} \setminus \{m\}}{\min} d(m', x_j) \right].$$
 (5)

The second term in both (4) and (5), namely  $\min_{m' \in \mathcal{M}_l} d(m', x_j)$  and  $\min_{m' \in \mathcal{M} \setminus \{m\}} d(m', x_j)$ , can be determined by caching the smallest and the second smallest distances from each point to the previous set of medoids, namely  $\mathcal{M}_l$  in (4) and  $\mathcal{M}$  in (5). Therefore, in both (4) and (5), we only need to compute the distance once for each summand. As a result, PAM needs  $O(kn^2)$  distance computations for the k greedy searches in the entire BUILD step and  $O(kn^2)$  distance computations for each SWAP iteration.

#### 143 **Bandit-PAM**

At the core of the PAM algorithm is the  $O(n^2)$  BUILD search (4), which is repeated k times for initialization, and the  $O(kn^2)$  SWAP search (5), which is repeated until convergence. We first show that both searches share a similar mathematical structure, and then show such a structure can be optimized efficiently using a bandit-based randomized algorithm, thus giving rise to Bandit-PAM. Rewriting the BUILD search (4) and the SWAP search (5) in terms of the change in total loss yields

BUILD: 
$$\underset{x \in \mathcal{X} \setminus \mathcal{M}_{l}}{\operatorname{arg \, min}} \frac{1}{n} \sum_{j=1}^{n} \left[ \left( d(x, x_{j}) - \min_{m' \in \mathcal{M}_{l}} d(m', x_{j}) \right) \wedge 0 \right],$$
 (6)

SWAP: 
$$\underset{(m,x)\in\mathcal{M}\times(\mathcal{X}\setminus\mathcal{M})}{\arg\min} \frac{1}{n} \sum_{j=1}^{n} \left[ \left( d(x,x_j) - \min_{m'\in\mathcal{M}\setminus\{m\}} d(m',x_j) \right) \wedge 0 \right]. \tag{7}$$

One may notice that the above two problems share the following similarities. First, both are searching over a finite set of parameters: n-l points in the BUILD search and k(n-k) swaps in the SWAP search. Second, both objective functions have the form of an average of an O(1) function evaluated over a finite set of reference points. We formally describe the shared structure:

Shared Problem: 
$$\underset{x \in \mathcal{S}_{tar}}{\operatorname{arg \, min}} \frac{1}{|\mathcal{S}_{ref}|} \sum_{x_j \in \mathcal{S}_{ref}} g_x(x_j),$$
 (8)

for target points  $S_{\text{tar}}$ , reference points  $S_{\text{ref}}$ , and an objective function  $g_x(\cdot)$  that dependents on the target point x. Then both the BUILD search and the SWAP search can be written as instances of Problem (8) with:

BUILD: 
$$S_{tar} = \mathcal{X} \setminus \mathcal{M}_l$$
,  $S_{ref} = \mathcal{X}$ ,  $g_x(x_j) = \left(d(x, x_j) - \min_{m' \in \mathcal{M}_l} d(m', x_j)\right) \wedge 0$ , (9)

SWAP: 
$$S_{tar} = \mathcal{M} \times (\mathcal{X} \setminus \mathcal{M}), \ S_{ref} = \mathcal{X}, \ g_x(x_j) = \left(d(x, x_j) - \min_{m' \in \mathcal{M} \setminus \{m\}} d(m', x_j)\right) \wedge 0.$$
 (10)

Crucially, in the SWAP search, each *pair* of medoid-and-non-medoid points (m, x) is treated as one target point in  $S_{tar}$  in this new formulation.

#### 3.1 Adaptive Search for the Shared Problem

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Recall that the computation of  $g(x_j)$  is O(1). A naive, explicit method would require  $O(|\mathcal{S}_{tar}||\mathcal{S}_{ref}|)$  computations of  $g(x_j)$  to solve Problem (8). However, as shown in previous works [2, 3], a randomized search would return the correct result with high confidence in  $O(|\mathcal{S}_{tar}|\log |\mathcal{S}_{ref}|)$  computations

## **Algorithm 1** Adaptive-Search $(S_{tar}, S_{ref}, g_x(\cdot), batchsize, \delta, \sigma_x)$

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1: \mathcal{S}_{solution} \leftarrow \mathcal{S}_{tar}
                                                                                                              ▷ Set of potential solutions to Problem (8)
                                                                                                                   > Number of reference points evaluated
 2: n_{\text{used\_ref}} \leftarrow 0
 3: For all x \in \mathcal{S}_{tar}, set \hat{\mu}_x \leftarrow 0, C_x \leftarrow \infty
                                                                                             ▶ Initial mean and confidence interval for each arm
 4: while n_{\text{used\_ref}} < |\mathcal{S}_{\text{ref}}| and |\mathcal{S}_{\text{solution}}| > 1 do
              Draw a batch of batchsize samples with replacement from reference S_{\text{ref batch}} \subset S_{\text{ref}}
              for all x \in \mathcal{S}_{\text{solution}} do \hat{\mu}_x \leftarrow \frac{n_{\text{used},\text{ref}}\hat{\mu}_x + \sum_{y \in \mathcal{S}_{\text{ref}} \text{ batch}} g_x(y)}{n_{\text{used},\text{ref}} + \text{batchsize}}
C_x \leftarrow \sigma_x \sqrt{\frac{2\log(\frac{1}{\delta})}{n_{\text{used},\text{ref}} + \text{batchsize}}}
 6:
 7:
                                                                                                                                                  8:
                                                                                                                                       ▶ Update confidence interval
              S_{\text{solution}} \leftarrow \{x: \hat{\mu}_x - C_x \leq \min_y (\hat{\mu}_y + C_y)\} \triangleright \text{Remove points that can no longer be solution}
 9:
10:
              n_{\text{used\_ref}} \leftarrow n_{\text{used\_ref}} + \text{batchsize}
11: if |S_{\text{solution}}| = 1 then
12:
               return x^* \in \mathcal{S}_{\text{solution}}
13: else
14:
              Compute \mu_x exactly for all x \in \mathcal{S}_{\text{solution}}
              return x^* = \arg\min_{x \in \mathcal{S}_{\text{solution}}} \mu_x
15:
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of  $g(x_j)$ . Specifically, for each target x in Problem (8), let  $\mu_x = \frac{1}{|\mathcal{S}_{\text{ref}}|} \sum_{x_j \in \mathcal{S}_{\text{ref}}} g_x(x_j)$  denote its objective function. Computing  $\mu_x$  exactly takes  $O(|\mathcal{S}_{\text{ref}}|)$  computations of  $g(x_j)$ , but we can instead estimate  $\mu_x$  with fewer computations by drawing  $J_1, J_2, ..., J_{n'}$  independent samples uniformly with replacement from  $[|\mathcal{S}_{\text{ref}}|]$ . Then,  $E[g(x_{J_i})] = \mu_x$  and  $\mu_x$  can be estimated as  $\hat{\mu}_x = \frac{1}{n'} \sum_{i=1}^{n'} g(x_{J_i})$ , where n' determines the estimation accuracy. To estimate the solution to Problem (8) with high confidence, we can then choose to sample different targets in  $\mathcal{S}_{\text{tar}}$  to different degrees of accuracy. Intuitively, promising targets with small values of  $\mu_x$  should be estimated with high accuracy, while less promising ones can be discarded without being evaluated on too many reference points.

The specific adaptive estimation procedure is described in Algorithm 1. It can be viewed as a batch version of the conventional UCB algorithm [21, 39] and is easier to implement. The algorithm uses the set  $\mathcal{S}_{\text{solution}}$  to track all potential solutions to Problem (8);  $\mathcal{S}_{\text{solution}}$  is initialized as the set of all target points  $\mathcal{S}_{\text{tar}}$ . For each potential solution  $x \in \mathcal{S}_{\text{solution}}$ , the algorithm maintains its mean objective estimate  $\hat{\mu}_x$  as well as a confidence interval  $C_x$ , where the latter depends on the exclusion probability  $\delta$  as well as the dispersion parameter  $\sigma_x$ .

In each iteration, a new batch of reference points  $S_{\text{ref\_batch}}$  is evaluated for all potential solutions in 176  $S_{\text{solution}}$ , making the estimate of  $\hat{\mu}_x$  more accurate. Based on the current estimate, if a target's lower 177 confidence bound  $\hat{\mu}_x - C_x$  is still greater than the upper confidence bound of the most promising 178 target  $\min_y (\hat{\mu}_y + C_y)$ , we remove it from the set of possible solutions  $\mathcal{S}_{\text{solution}}$ . This process continues 179 until there is only one point in  $S_{\text{solution}}$  or until we have sampled more reference points than in the 180 whole reference set. In the latter case, we know that the difference between the remaining targets 181 in  $S_{\text{solution}}$  is so subtle that an exact computation is more efficient. We then compute those targets' 182 objectives exactly and return the best target in the set. 183

#### 3.2 Algorithmic details

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Estimation of each  $\sigma_x$ : Bandit-PAM uses Algorithm 1 in both the BUILD step and each SWAP iteration, with input parameters specified in (9) and (10). In practice,  $\sigma_x$  is not known *a priori* and we estimate  $\sigma_x$  for each  $x \in |\mathcal{S}_{tar}|$  from the data. In the first batch of sampled reference points in Algorithm 1, we estimate each  $\sigma_x$  as:

$$\sigma_x = \mathrm{STD}_{y \in \mathcal{S}_{\text{ref batch}}} g_x(y) \tag{11}$$

where STD denotes standard deviation. Intuitively, this allows for smaller confidence intervals in later iterations, especially in the BUILD step, when we expect the average arm returns to become smaller as we add more medoids (since we are taking the minimum over a larger set on the RHS of Eq. (4)). We also allow for arm-dependent  $\sigma_x$ , as opposed to a fixed global  $\sigma$ , which allows for narrower confidence intervals for arms whose returns are heavily concentrated (e.g., distant outliers). Empirically, this results in significant speedups and results in fewer arms being computed exactly

(Line 14 in Algorithm 1). In all experiments, the batchsize is set to 100 and the error probability  $\delta$  is set to  $\delta = \frac{1}{1000|S_{tar}|}$  in Algorithm 1. Empirically, this value of batch size and this setting of  $\delta$  are such that Bandit-PAM recovers the same results in PAM in almost all cases.

198 **Combination with FastPAM1:** We also combine Bandit-PAM with the FastPAM1 optimization 199 [35]. We discuss this optimization in Appendix 1.2.

## 4 Analysis of the Algorithm

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The goal of Bandit-PAM is to track the optimization trajectory of the standard PAM algorithm, ultimately identifying the same set of k medoids with high probability. In this section, we formalize this statement and provide bounds on the number of distance computations required by Bandit-PAM.

We will assume that both PAM and Bandit-PAM place a hard constraint T on the maximum number of SWAP steps that are allowed. Notice that, as long as Bandit-PAM finds the correct solution to the search problem (6) at each BUILD step and to the search problem (7) at each SWAP step, it will reproduce the sequence of BUILD and SWAP steps of PAM identically, returning the same set of t medoids in the end. The hard constraint t guarantees that, even if the trajectories of PAM and Bandit-PAM deviate from each other, at most t t calls to Algorithm 1 will be performed.

Consider one such call to Algorithm 1 and suppose  $x^*$  is the optimal target point (i.e., the one with minimum  $\mu_x$ ). For another target point  $x \in \mathcal{S}_{tar}$ , let  $\Delta_x = \mu_x - \mu_{x^*}$ . To state the next result, we will assume that, for a randomly sampled reference point, say  $x_J$ , the random variable  $Y = g_x(x_J)$  is  $\sigma$ -sub-Gaussian; i.e., that  $\Pr(|Y - E[Y]| > t) < 2 \exp\left(-t^2/(2\sigma^2)\right)$ , for some known parameter  $\sigma$ . In addition, we assume that the data is generated in such a distribution that the mean rewarded  $\mu_i$ 's follow a sub-Gaussian distribution (see Sec. 6 for a discussion).

Theorem 1. If Bandit-PAM is run on a dataset  $\mathcal{X}$  with  $\delta=n^{-3}$ , then it returns the same set of k medoids as PAM with probability 1-2(k+T)/n. Furthermore, the total number of distance computations  $M_{\mathrm{total}}$  required satisfies

$$E[M_{\text{total}}] = O((k+T)n\log n).$$

When the number of desired medoids k is a constant and the number of allowed SWAP steps is small (which is often sufficient in practice as discussed in Sec. 6). Theorem 1 implies that only  $O(n \log n)$  distance computations are necessary to reproduce the results of PAM with high probability.

In order to prove Theorem 1, we prove a more detailed result for each call that Bandit-PAM makes to Algorithm 1. For this more specific case, we assume that, for target point x,  $g_x(x_J)$  is  $\sigma_x$ -sub-Gaussian, where  $\sigma_x$  is a parameter specific to x (and can change across different calls to Algorithm 1). As it turns out, in practice one can estimate each  $\sigma_x$  by performing a small number of distance computations. Allowing  $\sigma_x$  to be estimated separately for each arm is beneficial in practice, as discussed in Sec. 6. The following theorem is proved in Appendix 3.

Theorem 2. For  $\delta = n^{-3}$ , with probability at least  $1 - \frac{2}{n}$ , Algorithm 1 returns the correct solution to (6) (for a BUILD step) or (7) (for a SWAP step), using a total of M distance computations, where

$$E[M] \le 4n + \sum_{x \in \mathcal{X}} \min \left[ \frac{24}{\Delta_x^2} \left( \sigma_x + \sigma_{x^*} \right)^2 \log n + \text{batchsize}, 2n \right].$$

While the assumption that  $\sigma_x$  is known for every x may seem excessive, it is worth pointing out that Algorithm 1 does not need to know all  $\sigma_x$ s exactly and an upper bound is sufficient. Notice that, if a random variable is  $\sigma$ -sub-Gaussian, it is also  $\sigma'$ -sub-Gaussian for  $\sigma' > \sigma$ . Hence, if we have a universal upper bound  $\sigma_{\rm ub} > \sigma_x$  for all x, the algorithm can be run with  $\sigma_{\rm ub}$  replacing each  $\sigma_x$ . In that case, a direct consequence of Theorem 2 is that the total number of distance computations per call to Algorithm 1 satisfies

$$E[M] \le 4n + \sum_{x \in \mathcal{X}} 96 \frac{\sigma_{\text{ub}}^2}{\Delta_x^2} \log n + \text{batchsize} \le 4n + 96 \left(\frac{\sigma_{\text{ub}}}{\min_x \Delta_x}\right)^2 n \log n.$$
 (12)

Furthermore, as proved in Appendix 2 of Bagaria et al. [2], such an instance-wise bound converts to an  $O(n \log n)$  bound when  $\mu_i$ 's follow a Sub-Gaussan distribution. Moreover, from Theorem 2, the

Table 1: Description of Datasets

Name	Size	Distance Metric	Dimensionality
MNIST scRNA-seq HOC4	70000 40000 3,360	$l_2$ , cosine distance $l_1$ Tree edit distance	784 10170 ∞

probability that Algorithm 1 does not return the target point x with the smallest value of  $\mu_x$  is at most 2/n. By the union bound, the probability that Bandit-PAM does not return the same set of k medoids as PAM is at most 2(k+T)/n. Moreover, since at most k+T calls to Algorithm 1 are made, from (12) we see that the total number of distance computations  $M_{\text{total}}$  required by Bandit-PAM satisfies  $E[M_{\text{total}}] = O((k+T)n\log n)$ . This proves Theorem 1.

## **5 Empirical Results**

We run experiments on three real-world datasets to validate the expected behavior of Bandit-PAM: the MNIST hand-written digits dataset [22], the 10x Genomics 68k PBMCs scRNA-seq dataset [40], and the Code.org Hour Of Code #4 (HOC4) coding exercise submission dataset (code.org), all of which are publicly available.

**Datasets.** The MNIST dataset [22] consists of 70,000 black-and-white images of handwritten digits, where each digit is represented as a 784 dimensional vector. We consider two distance metrics, namely  $l_2$  distance and cosine distance. The scRNA-seq dataset contains the gene expression levels of 10,170 different genes in each of 40,000 cells after standard filtering. We consider  $l_1$  distance that is popular for clustering scRNA-seq data [31] has often processed scRNA-seq datasets with  $l_1$  distance. The HOC4 dataset from Code.org [8] consists of 3,360 unique solutions to a block-based programming exercise on Code.org. Solutions to the programming exercise are represented as abstract syntax trees (ASTs), and we consider the tree edit distance to quantify the similar between solutions. See Table 1 for a summary.

**Setup.** In Subsec. 5.1, we show that Bandit-PAM returns the same results as PAM. We also compare the clustering results with other popular k-medoids clustering algorithms in terms of the clustering loss (3), including FastPAM [35], CLARANS [30], and Voronoi Iteration [33]. In Subsec. 5.2, we demonstrate that Bandit-PAM scales linearly in the number of samples n for all datasets and all metrics considered. Each parameter setting was repeated 10 times with data subsampled from the original dataset. 95% confidence intervals are provided.

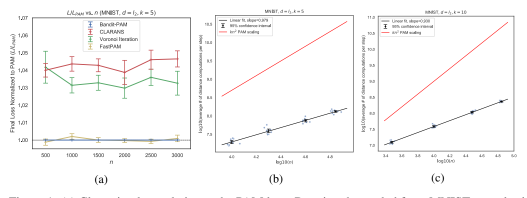


Figure 1: (a) Clustering loss relative to the PAM loss. Data is subsampled from MNIST, sample size n varies from 500 to 3000, k=5 and 95% confidence intervals are provided. Bandit-PAM always returns the same solution as PAM and hence has loss ratio 1. FastPAM has a comparable performance, while the other two algorithms are significantly worse. (b-c) Average number of distance calls per iteration vs sample size n for MNIST and  $l_2$  distance with (b) k=5 and (c) k=10. The plot is shown on a log-log scale. Lines of best fit (black) are plotted, as are reference lines demonstrating the expected scaling of PAM (red).

#### 5.1 Clustering/loss quality

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Figure 1 (a) shows the relative losses of algorithms with respect to the loss of PAM. Bandit-PAM and three other baselines, namely FastPAM [35], CLARANS [30], and Voronoi Iteration [33], are considered. We note that FastPAM is different from FastPAM1 mentioned before; it takes  $O(n^2)$  for each SWAP step but does not guarantee the same solution as PAM. Bandit-PAM always returns the same solution as PAM and hence has loss ratio 1. FastPAM has a comparable performance, while the other two algorithms are significantly worse.

#### Scaling with n for different datasets, distance metric, and k values 5.2

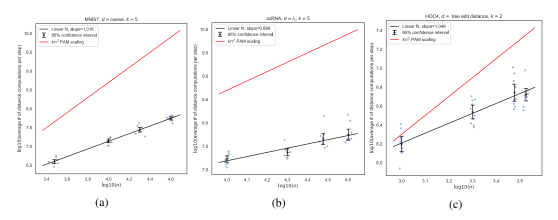


Figure 2: Average number of distance calls per iteration vs sample size n, for (a) MNIST and cosine distance, (b) scRNA-seq and  $l_1$  distance, and (c) HOC4 and tree edit distance. The plot is shown on a log-log scale. Lines of best fit (black) are plotted, as are reference lines demonstrating the expected scaling of PAM (red).

We next consider the number of distance calls per iteration as the sample size increases. The number 271 of distance calls per iteration is defined as the total distance calls divided by the number of SWAP steps plus 1, where the 1 corresponds to the BUILD step. We choose to look at this quantity to account for different number of SWAPs for different runs, in order to provide a fair comparison.

If the complexity is linear, then the slope would be 1 in the log-log plot. Indeed, as shown in Figure 275 1 (b-c), the slope for k = 5 and k = 10 are 0.979 and 0.930, respectively, indicating the scaling is 276 linear in n for different values of k. 277

In addition, as shown in Figure 2, the slopes of the log-log plot are 1.018, 0.899 and 1.046 for 278 MNIST with cosine distance, scRNA-seq with  $l_1$  distance, HOC4 with tree edit distance, respectively, 279 validating our theory that Bandit-PAM takes almost linear number of distance evaluations per iteration 280 for different datasets and different distance metrics. 281

#### **Discussion**

In all experiments, we have observed that the numbers of SWAPs are very small, typically fewer than 10, justifying the assumption of having an upper limit on the PAM SWAP step prior to running the algorithm in Sec. 4.

We observe that for all datasets, the randomly sampled distances have an empirical distribution 286 similar to Gaussian distribution (Appendix Figures 4-5), justifying the Sub-Gaussian assumption in 287 Sec. 4. In addition, we observe that the Sub-Gaussian parameters are different for different steps 288 and different points (Appendix Figures 2), justifying the adaptive estimation of the sub-Gaussianity 289 parameters in SubSec. 3.2. 290

In addition, the distribution of the true arm parameters also mostly do have a heavy-tailed distribution 291 (Appendix Figure 3), justifying the distributional assumption of  $\mu_i$ 's in Sec. 4.

## Broader Impact

In this work, we proposed an algorithm that accelerated finding solutions to the k-medoids problem 294 295 while producing comparable – and usually equivalent – final cluster assignments. Our work enables the discovery of high-quality medoid assignments in very large datasets, including some on which 296 prior algorithms were prohibitively expensive. A potential negative consequence of this is that 297 practitioners may be incentivized to gather and store larger amounts of data now that it can be 298 meaningfully processed, in a phenomenon more generally described as induced demand [11]. This 299 incentive realignment could potentially result in negative externalities such as an increase in energy 300 consumption and carbon footprints. 301

We also anticipate, however, that Bandit-PAM will enable several beneficial applications in 302 biomedicine, education, and fairness. For example, the evolutionary pathways of infectious diseases 303 could possibly be constructed from the medoids of genetic sequences available at a given point in 304 time, if prior temporal information about these sequences' histories is not available. Similarly, the 305 medoids of patients infected in a disease outbreak may elucidate the origins of outbreaks, as did 306 prior analyses of cholera outbreaks using Voronoi Iteration [6]. Our application to the HOC4 dataset 307 also suggests a method for scaling personalized feedback to individual students in online courses. If 308 limited resources are available, instructors can choose to provide feedback on just the *medoids* of 309 submitted solutions instead of exhaustively providing feedback on every unique solution, of which 310 there may be several thousand. Instructors can then refer individual students to the feedback provided 311 for their closest medoid. We anticipate that this approach can be applied generally for students of 312 Massive Open Online Courses (MOOCs), thereby enabling more equitable access to education and 313 personalized feedback for students.. In particular, especially with recent interest in online learning, 314 we hope that our work will improve the quality of learning for students worldwide.

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