

# Scaling Density-Based Clustering to Large Collections of Sets

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## ABSTRACT

We study techniques for clustering large collections of sets into DBSCAN clusters. Sets are often used as a representation of complex objects to assess their similarity. The similarity of two objects is then computed based on the overlap of their set representations, for example, using Hamming distance. Clustering large collections of sets is challenging. A baseline that executes the standard DBSCAN algorithm suffers from poor performance due to the unfavorable neighborhood-by-neighborhood order in which the sets are retrieved. The DBSCAN order requires the use of a symmetric index, which is less effective than its asymmetric counterpart. Precomputing and materializing the neighborhoods to gain control over the retrieval order often turns out to be infeasible due to excessive memory requirements.

We propose a new, density-based clustering algorithm that processes data points in any user-defined order and does not need to materialize neighborhoods. Instead, so-called backlinks are introduced that are sufficient to achieve a correct clustering. Backlinks require only linear space while there can be a quadratic number of neighbors. To the best of our knowledge, this is the first DBSCAN-compliant algorithm that can leverage asymmetric indexes in linear space. Our empirical evaluation suggests that our algorithm combines the best of two worlds: it achieves the runtime performance of materialization-based approaches while retaining the memory efficiency of non-materializing techniques.

## 1 INTRODUCTION

We consider the problem of partitioning large collections of sets into DBSCAN [15] clusters. Our work is motivated by a process mining use case at Celonis SE that models processes as sets. A *process* is a sequence of timestamped activities. Large companies store hundreds of millions of activities in millions of processes. In order to analyze the processes, they should be clustered into groups of similar activity sequences that can be further explored [5, 22, 39]. To this end, a process is represented by the set of all its neighboring activity pairs, e.g., the process with the activity sequence  $(S, O, P, H, R, F, E)$  (Start, Order, Pay, sHip, Return good, reFund, End) is represented by the set  $\{(S, O), (O, P), (P, H), (H, R), (R, F), (F, E)\}$ . The similarity of two processes is then assessed by the Hamming distance<sup>1</sup> of their set representations.

Sets are used in many other applications [29] to represent objects for the purpose of clustering, e.g., sales may be represented by sets of product categories, photos by sets of tags and title words, user interactions on a website by sets of visited links, users of a social network by their group memberships, or users of a music streaming platform by sets of tracks they listen to.

<sup>1</sup>Hamming distance  $H(r, s) = |r \cup s| - |r \cap s|$  for two sets  $r$  and  $s$ .

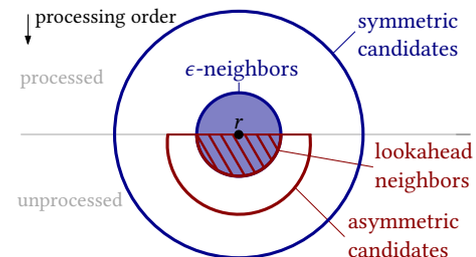


Figure 1: Symmetric candidates with  $\epsilon$ -neighbors (blue); asymmetric candidates with lookahead neighbors (red).

The popular DBSCAN (Density-Based Spatial Clustering of Applications with Noise) algorithm [15] identifies clusters of arbitrary shape without requiring the number of clusters as input. Intuitively, DBSCAN finds dense regions that are separated by regions of lower density. The density of a region (given a distance function between pairs of data points) is defined by two parameters: the number of neighbors, *minPts*, and the radius,  $\epsilon$ , of the neighborhood. A data point is called *core point* (i.e., it is at the core of a dense region) if it has at least *minPts* neighbors (including itself) within radius  $\epsilon$ ; a non-core point in the  $\epsilon$ -neighborhood of a core point is a *border point* (i.e., it is at the border of a dense region); all other points are *noise* [37].

The runtime of the DBSCAN algorithm heavily depends on the efficiency of the neighborhood computation. In our experiments, the neighborhood computation accounts for up to 99% of the overall runtime for some datasets. Therefore, in order to efficiently cluster large collections of sets, effective indexing techniques for sets are required.

Similarity indexes for sets have been proposed in the context of  $\epsilon$ -neighborhood joins, which are executed in an index nested loop fashion. A prominent representative is the *prefix index* [1, 8], which is linear in size and has been shown to be highly effective [29]. The *symmetric* prefix index returns the complete  $\epsilon$ -neighborhood for a given query point  $r$ . The *asymmetric* prefix index assumes a processing order on the sets in  $R$  and retrieves only the *lookahead neighbors*: the  $\epsilon$ -neighbors that follow  $r$  in processing order. A typical processing order for sets is based on the set sizes (ties broken arbitrarily). The *asymmetric* prefix index further leverages the length information to avoid many of the candidates that the symmetric index must inspect (among the unprocessed sets). Figure 1 illustrates the  $\epsilon$ -neighborhood, the lookahead neighbors, and the candidate regions of symmetric and asymmetric index, respectively. The region above the gray line represents the sets that have been processed before  $r$ , the region below the gray line are unprocessed sets. The circles and semicircles show subset relationships.

Clearly, the asymmetric prefix index is preferable in terms of effectiveness over its symmetric counterpart. Unfortunately, there is an inherent mismatch between the asymmetric index and the DBSCAN algorithm. DBSCAN suffers from the following issues when executed with the asymmetric index: (1) *Core status*

*problem*: The lookahead neighbors of  $r$  are not sufficient to update the core status of  $r$ . (2) *Border vs. Noise problem*: To distinguish border points from noise, a border point must be visible from a core point, which is not guaranteed by the asymmetric lookahead neighborhood. (3) *Disconnected clusters*: To guarantee connected clusters, DBSCAN imposes a (partial) processing order on the neighborhood computations: all core points of the current cluster must be expanded (i.e., their neighborhood must be computed) before any point belonging to a different cluster is processed.

A well-known clustering approach [3] is based on a self-join that precomputes and materializes all neighborhoods. The pre-computed neighborhoods are then used while executing DBSCAN. This approach can leverage the asymmetric index and is efficient in runtime. Unfortunately, this join-based technique requires quadratic memory in the worst case and suffers from a large memory footprint in practice. For example, for our social media dataset (LIVEJ) that stores the interests of 3.1M users, this approach requires almost 100GB of memory.

Summarizing, applications that must cluster large collections of sets have two options, which we call Sym-Clust and Join-Clust. (1) Sym-Clust: Retrieve the full  $\epsilon$ -neighborhoods in the processing order imposed by DBSCAN using the symmetric index. (2) Join-Clust: Compute and materialize neighborhoods in a join using the effective asymmetric index. None of the options is satisfying: Sym-Clust runs almost up to an order of magnitude slower than Join-Clust, while Join-Clust is infeasible for many datasets and parameter settings due to its excessive memory usage.

We propose a new clustering algorithm, *Spread*, that computes correct DBSCAN clusters using the asymmetric prefix index. *Spread* runs in linear space and does not need to materialize the (quadratic-size) neighborhoods. *Spread* avoids symmetric neighbor computations, therefore reducing the number of neighbors retrieved by Sym-Clust. So-called *backlinks* are introduced to achieve a correct clustering. Backlinks are dynamically added and removed as required and occupy only a small fraction of the memory that is used by materialized neighborhoods. *Spread* maintains a graph of subclusters in a disjoint-set data structure and guarantees that connected components in the resulting graph represent correct DBSCAN clusters.

In general, *Spread* can process data points in any *user-defined order* given an index that retrieves the lookahead neighbors, i.e., all data points that follow the query point in the user-defined processing order. In our usage scenario – set clustering – the processing order is defined by the set sizes (ties broken arbitrarily) and the asymmetric prefix index retrieves lookahead neighbors.

Summarizing, our contributions are the following:

- We propose *Spread*, a novel algorithm for partitioning large collections of sets into DBSCAN clusters. To the best of our knowledge, this is the first linear space DBSCAN-compliant algorithm that leverages the *asymmetric* prefix index for sets.
- We introduce the new concept of *backlinks* that keep sufficient information to build correct clusters independently of the processing order that the user imposes on *Spread*. We prove invariants for backlinks and the correctness of our approach.
- Our extensive empirical evaluation on 13 real-world datasets suggests that *Spread* is as fast as Join-Clust (that materializes all neighborhoods) while being competitive in memory usage with Sym-Clust (that computes all neighborhoods on the fly).

The remainder of this paper is organized as follows. In Section 2, we cover the background on  $\epsilon$ -neighborhood and set similarity, indexing techniques for sets, and density-based clustering, and we define the *density-based set clustering problem*. Section 3 presents the two baseline approaches for density-based set clustering, Join-Clust and Sym-Clust. In Section 4, we present *Spread*, our time- and space-efficient solution for density-based set clustering. We evaluate our solution against the baseline algorithms and discuss the results in Section 5. Related work is summarized in Section 6. Finally, Section 7 concludes this paper.

## 2 BACKGROUND & PROBLEM DEFINITION

We revisit set similarity indexes and density-based clustering, and define our problem. To simplify the presentation, we focus on prefix indexes for the Hamming distance. Our results, however, extend to other distance and similarity measures (e.g., Jaccard or Cosine) and the respective indexes [12, 29, 40]. The required adaptations of the index that have been studied in the context of set similarity joins [29, 49] (e.g., prefix length, size lower bound, equivalent overlap) also apply to our scenario.

### 2.1 Set Similarity and $\epsilon$ -Neighborhood

$R$  is a collection of  $n$  unique sets, each set  $r \in R$  consists of unique tokens  $t_1, \dots, t_m$ ,  $|r| = m$ . The *processing order*,  $>$ , is a total order defined over  $R$ . The similarity between two sets  $r$  and  $s$  is assessed by the Hamming distance,  $H(r, s) = |r \cup s| - |r \cap s|$ , which counts the number of tokens that exist only in one of the sets, e.g.,  $H(r_1, r_2) = 4$  and  $H(r_2, r_3) = 3$  for the sets in Figure 2.

The  $\epsilon$ -neighborhood of set  $r$  includes  $r$  and all sets within distance  $\epsilon$  from  $r$ ,  $N_\epsilon(r) = \{s \in R \mid H(r, s) \leq \epsilon\}$ . A *region query* on  $r$  computes  $N_\epsilon(r)$ . A set  $r$  splits its  $\epsilon$ -neighborhood into two disjoint parts based on the processing order: the *lookahead neighbors* that follow  $r$  in processing order,  $N_\epsilon^>(r) = \{s \in N_\epsilon(r) \mid s > r\}$  and the *preceding neighbors*,  $N_\epsilon^<(r) = \{s \in N_\epsilon(r) \mid s < r\}$ .

### 2.2 Indexing Techniques for Sets

*Prefix Filter and Inverted Index*. A naive approach computes a region query  $N_\epsilon(r)$  by verifying the predicate  $H(r, s) \leq \epsilon$  for all sets  $s \in R$ . An effective indexing technique, which was originally developed for set similarity joins [2, 29], is based on the so-called prefix filter. The prefix,  $\pi_r$ , of a set  $r$  consists of the first  $\pi$  tokens of  $r$  according to some total token order (which must be the same for all sets). The prefix length depends on the distance function and is  $\pi = \epsilon + 1$  for the Hamming distance. Figure 2 shows the prefix of three sets for distance threshold  $\epsilon = 3$  and a numerical token order. The prefix filter works best if the tokens in the prefix are infrequent, thus the tokens are typically ordered by ascending global token frequency.

A set  $s \in R$  can be in the  $\epsilon$ -neighborhood  $N_\epsilon(r)$  only if the prefixes of  $r$  and  $s$  share at least one token, i.e.,  $H(r, s) \leq \epsilon \Rightarrow \pi_r \cap \pi_s \neq \emptyset$  (assuming  $|r| + |s| > \epsilon$ ; otherwise  $r$  and  $s$  are always similar). Therefore, if two sets do not share a token in the prefix, the pair can be safely pruned. If two sets  $r$  and  $s$  share a prefix token,  $(r, s)$  is a *candidate pair* and must undergo *verification*, i.e., the predicate  $H(r, s) \leq \epsilon$  must be evaluated. Candidates that fail verification are *false positives*. Mann et al. [29] discuss efficient prefix-based verification.

*Symmetric Prefix Index*. An inverted index on the prefix tokens is used to retrieve candidate pairs efficiently. The inverted index maps prefix tokens to sets that contain that token in the prefix. A lookup of set  $r$  retrieves all lists of the prefix tokens of  $r$ . The

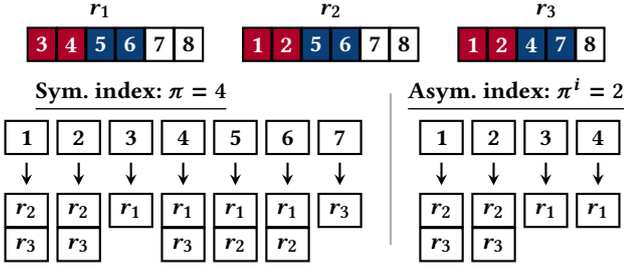


Figure 2: Symmetric and asymmetric prefix index,  $\epsilon = 3$ .

union of these lists (except  $r$  itself) are the candidates of  $r$ . The index is *symmetric* and returns the  $\epsilon$ -neighborhood of  $r$ .

For example, the candidates for  $r_2$  returned by the symmetric index,  $\pi = \epsilon + 1 = 4$ , in Figure 2 are  $\{r_1, r_3\}$  (resulting from the union of  $[r_2, r_3]$  for token 1,  $[r_2, r_3]$  for token 2,  $[r_1, r_2]$  for token 5, and  $[r_1, r_2]$  for token 6). Candidate  $r_1$  is a false positive since  $H(r_1, r_2) > \epsilon$ ;  $r_3$  is a true positive due to  $H(r_2, r_3) \leq \epsilon$ .

*Asymmetric Prefix Index.* We construct an *asymmetric* prefix index that returns only the lookahead neighbors,  $N_\epsilon^>(r)$ . To this end, we define a length-based processing order on  $R$  (longest to shortest):  $r$  precedes  $s$  if  $|r| > |s|$ , i.e.,  $|r| > |s| \Rightarrow s > r$ ; ties ( $|r| = |s|$ ) are broken by the lexicographic order of the sorted sets.

Since we are interested only in sets  $s \in N_\epsilon(r)$  that are no larger than  $r$ ,  $|s| \leq |r|$ , we need to index only a subset of the prefix tokens: the tokens in the so-called *indexing prefix* [49]. The so-called *probing prefix* of the lookup set,  $r$ , remains of length  $\pi = \epsilon + 1$ . For the Hamming distance, the indexing prefix is of length  $\pi^i = \lfloor \frac{\pi}{2} + 1 \rfloor$ . For  $\epsilon > 0$ , the probing prefix is always longer than the indexing prefix, e.g.,  $\pi = 4$  and  $\pi^i = 2$  for  $\epsilon = 3$ . A shorter prefix results in fewer candidates and renders the asymmetric index more effective than its symmetric counterpart.

In the case of  $r_2$ , the asymmetric index,  $\pi^i = 2$ , in Figure 2 returns only a true positive candidate,  $r_3$ . The false positive candidate,  $r_1$ , which is returned by the symmetric index, is avoided.

## 2.3 Density-Based Clustering

We formally define DBSCAN clusters and the related concepts. A set  $r$  represents a point to be clustered. The *density* of  $r$  is the number of  $\epsilon$ -neighbors  $|N_\epsilon(r)|$  (cf. Section 2.1).

*Core, Border, Noise.* A set  $r$  is a *core point* iff the  $\epsilon$ -neighborhood of  $r$  contains at least  $\text{minPts}$  sets:  $r$  is core  $\Leftrightarrow |N_\epsilon(r)| \geq \text{minPts}$ . A set  $s$  is a *border point* iff it is in the  $\epsilon$ -neighborhood of a core point  $r$  and  $s$  is not core:  $s$  is border  $\Leftrightarrow s \in N_\epsilon(r) \wedge |N_\epsilon(r)| \geq \text{minPts} \wedge |N_\epsilon(s)| < \text{minPts}$ . All remaining sets in  $R$  are *noise*. We denote the set of core and border points with  $\mathcal{C}$  and  $\mathcal{B}$ , respectively. The set of noise points is  $\mathcal{N} = R \setminus (\mathcal{C} \cup \mathcal{B})$ .

*Density-Reachability.* Let  $r, s \in R$  and  $r$  is core:  $s$  is *directly density-reachable* from  $r$  iff  $s$  is in the  $\epsilon$ -neighborhood of  $r$ :  $r \blacktriangleright s \Leftrightarrow s \in N_\epsilon(r)$ . If there is a sequence of sets  $r_1, r_2, \dots, r_k$  with  $r_1 = r$  and  $r_k = s$ ,  $r_i \blacktriangleright r_{i+1}$  for  $1 \leq i < k$ ,  $s$  is *density-reachable* from  $r$ , denoted  $r \blacktriangleright \dots \blacktriangleright s$ . Two sets  $r, s$  are *density-connected* if there is a set  $x$  s.t. both  $r$  and  $s$  are density-reachable from  $x$ .

A *density-based cluster* is a subset  $C_i \subseteq R$  that satisfies two criteria [38]:

- (1) *Maximality*  $\mathbb{M}$ : For any two sets  $r, s \in R$ ,  $r \in C_i$ . If  $s$  is density-reachable from  $r$ , then  $s \in C_i$ . Formally,

$$\forall r, s \in R : r \in C_i \wedge r \blacktriangleright \dots \blacktriangleright s \Rightarrow s \in C_i$$

Table 1: Notation overview.

Notation	Description
$R$	a collection of sets
$r, s, x$	sets of $R$
$ r $	cardinality of set $r$
$r < s, r > s$	$r$ precedes/succeeds $s$ (in $R$ )
$H(r, s)$	the Hamming distance of two sets $r, s$
$\pi, \pi^i$	probing/indexing prefix
$\epsilon$	distance threshold
$\text{minPts}$	minimum density s.t. a set $r$ is core
$N_\epsilon(r)$	full $\epsilon$ -neighborhood of $r$
$N_\epsilon^<(r), N_\epsilon^>(r)$	preceding/lookahead neighbors of $r$
$r \blacktriangleright s$	$s$ is directly density-reachable from $r$
$r \blacktriangleright \dots \blacktriangleright s$	$s$ is density-reachable from $r$
$\mathcal{C}, \mathcal{B}, \mathcal{N}$	the set of core, border, and noise sets
$C_i$	a density-based cluster with id $i$

- (2) *Connectivity*  $\mathbb{C}$ : For any two sets  $r, s$  in  $C_i$ , there is a set  $x$  that density-connects  $r$  and  $s$ . Formally,

$$\forall r, s \in R : r, s \in C_i \Rightarrow \exists x \in C_i : r \blacktriangleleft \dots \blacktriangleleft x \blacktriangleright \dots \blacktriangleright s$$

*DBSCAN Clustering.* A border point may be part of multiple density-based clusters such that the clusters overlap. We define the *DBSCAN clustering* that partitions the data into non-overlapping clusters. The standard DBSCAN algorithm [15] produces a DBSCAN clustering.

*Definition 2.1.* Let  $R^* = R \setminus \mathcal{N}$  and  $C_1, C_2, \dots, C_k$  be density-based clusters such that  $\bigcup_{i=1}^k C_i = R^*$ . A DBSCAN clustering is a partitioning  $\Gamma = \{C'_1, C'_2, \dots, C'_k\}$ ,  $C'_i \subseteq C_i$ , such that  $\bigcup_{i=1}^k C'_i = R^*$ ,  $C'_i \cap C'_j = \emptyset$  for  $i \neq j$ .

A *subclustering* of a cluster  $C_i$ ,  $\psi_i = \{c_1, c_2, \dots, c_l\}$ , is a partitioning of  $C_i$  into  $1 \leq l \leq |C_i|$  non-empty, disjoint subclusters,  $c_j \subseteq C_i$ , such that  $\bigcup_{j=1}^l c_j = C_i$ ,  $c_j \cap c_k = \emptyset$  for  $j \neq k$ .

A *subcluster graph* of  $R^*$  is an undirected graph in which nodes are subclusters and an edge between two nodes can only exist if the respective nodes are in the same DBSCAN cluster.

## 2.4 The DBSCAN Algorithm

The standard DBSCAN algorithm [15] forms clusters by repeatedly picking a seed point from the set of unvisited data points (initially all points are unvisited). If the seed is a core point, it forms a new cluster with all points that are density-reachable from the seed and are not yet assigned to a cluster. The set of density-reachable points is computed by recursively adding the  $\epsilon$ -neighbors of all core points to the current cluster. The algorithm terminates when all points have been visited. Points that cannot be assigned to a cluster are noise.

## 2.5 Problem Statement

*Definition 2.2 (Density-Based Set Clustering).* Given a collection of sets  $R$ , a distance threshold  $\epsilon$ , and the neighborhood density  $\text{minPts}$ , the goal is to find a DBSCAN clustering  $\Gamma = \{C_1, C_2, \dots, C_k\}$  of  $R$ .

For sets, asymmetric indexes with a lookahead neighbor function  $N_\epsilon^>(r)$  promise the best performance (cf. Section 2.2). Given an ordering  $>$  on  $R$ , we strive for a time- and space-efficient algorithm that solves the density-based set clustering problem with an asymmetric index.

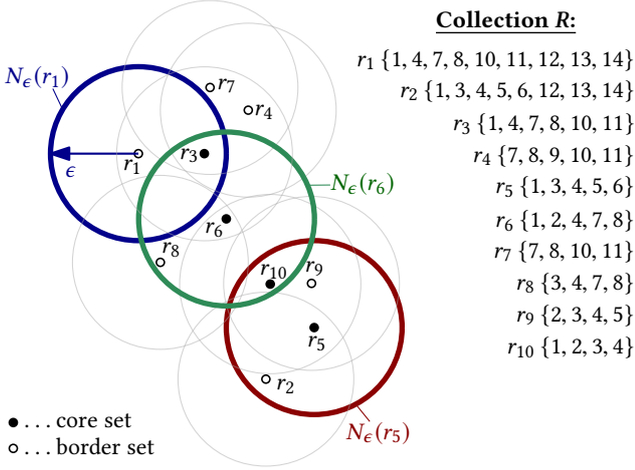


Figure 3: Running example,  $\epsilon = 3$ ,  $\text{minPts} = 4$ .

*Running Example.* Figure 3 shows an example collection  $R$  of ten sets,  $r_1$ – $r_{10}$ , and their neighborhoods for Hamming distance  $\epsilon = 3$ . Sets  $r_3$ ,  $r_5$ ,  $r_6$ , and  $r_{10}$ , are core sets; all sets  $r_1$ – $r_{10}$  form a single cluster.

### 3 BASELINE APPROACHES

This section presents two baseline solutions for the density-based set clustering problem. (1) *Sym-Clust* is memory-efficient and follows the standard DBSCAN approach with the symmetric prefix index to answer neighborhood queries on the fly. (2) *Join-Clust* is speed-optimized and materializes all  $\epsilon$ -neighborhoods using a state-of-the-art set similarity join algorithm [2] (which leverages the asymmetric prefix index) before the standard DBSCAN algorithm is executed.

Both baselines leverage state-of-the-art set indexes. We are not aware of other previous solutions that can outperform *Sym-Clust* or *Join-Clust* for the density-based set clustering problem. Note that using the standard DBSCAN [15] (rather than some advanced techniques presented in later works, cf. Section 6) is not a limiting factor: Most of the overall execution time is spent computing the neighborhoods, and prefix-based indexes are highly efficient in combination with efficient verification [29].

#### 3.1 Sym-Clust: DBSCAN with Inverted Index

When the standard DBSCAN algorithm (cf. Section 2.4) picks a seed point that is core, it forms a cluster with all points that are density-reachable from the seed. The density-reachable points are computed by pushing all core neighbors of the seed onto a stack. Then, each point on the stack is processed in the same manner (i.e., all its core neighbors are pushed onto the stack) until the stack is empty. All neighbors of core points retrieved in this process belong to the cluster.

The neighborhood queries will overlap to some extent. Assume  $r$  is processed before  $s$ ,  $s \in N_\epsilon(r)$ , then  $|N_\epsilon(s) \cap N_\epsilon(r)| \geq 2$  (at least  $r$  and  $s$  are in both neighborhoods). Since  $r$  assigns all its neighbors to the current cluster, only the non-overlapping neighbors of  $s$ ,  $N_\epsilon(s) \setminus N_\epsilon(r)$ , will further increase the cluster.

Figure 4 illustrates this observation for the neighborhoods of two example points  $r$  (black circle) and  $s$  (red circle): only the new, non-overlapping area of  $N_\epsilon(s)$  (shaded in red) is relevant for expanding the cluster.

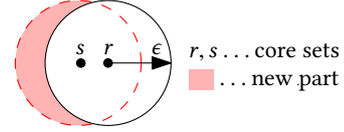


Figure 4: Redundant neighborhood queries.

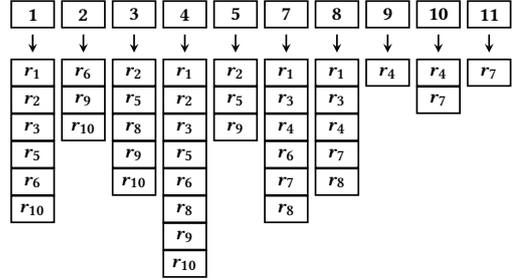


Figure 5: Symmetric prefix index on  $r_1$ – $r_{10}$ ,  $\epsilon = 3$ ,  $\pi = 4$ .

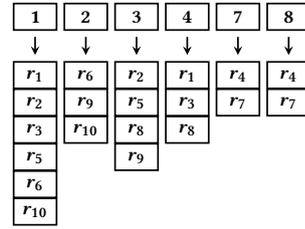


Figure 6: Asymmetric prefix index on  $r_1$ – $r_{10}$ ,  $\epsilon = 3$ ,  $\pi^i = 2$ .

The standard DBSCAN algorithm requires the use of a symmetric index since it assumes to see all neighbors of a point  $s$  when  $s$  is processed. The asymmetric index is not compatible with the standard DBSCAN algorithm: We cannot impose an order on the points such that all non-overlapping neighbors of  $s$  follow  $s$  in the processing order. Further, the size of the neighborhood of  $s$ ,  $|N_\epsilon(s)|$ , is required to decide its core status.

Figure 5 shows the symmetric prefix index for our running example,  $\epsilon = 3$ ,  $\pi = \epsilon + 1 = 4$ . We probe  $r_8 = \{3, 4, 7, 8\}$ . The prefix of  $r_8$  consists of all tokens in  $r_8$  (due to  $|r_8| = \pi$ ). The union of the respective index lists yields the candidates  $\{r_2, r_5, r_9, r_{10}, r_1, r_3, r_6, r_4, r_7\}$ . Note that the candidates include both sets that are smaller and sets that are larger than  $r_8$ .

The so-called *length filter* [2], an optimization of the symmetric prefix index that also applies to its asymmetric counterpart, prunes candidates  $r_2$  and  $r_1$ . Due to their length difference to  $r_8$ , they cannot be in the  $\epsilon$ -neighborhood of  $r_8$ . By ordering the lists in processing order (i.e., longer sets precede shorter sets, as illustrated in Figure 5), the length filter can prune the head (sets that are too long) and the tail (sets that are too short) of a list without inspecting all elements in head and tail, respectively.

All candidates that are not pruned by the length filter must undergo verification. Only  $r_6$  passes verification, therefore  $N_\epsilon(r_8) = \{r_8, r_6\}$ , and  $r_8$  is classified as non-core ( $\text{minPts} = 4$ ).

*Complexity Analysis.* We probe each set  $r \in R$  against the index once. With cost  $C$  for an index lookup and  $n = |R|$ , the runtime is  $O(n \cdot C)$ ;  $C = O(n)$  since  $r$  may have  $O(n)$  neighbors, thus the overall runtime of *Sym-Clust* is  $O(n^2)$ . The symmetric index is of linear size leading to space complexity  $O(n)$ .

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**Algorithm 1:** Materialize-Neighborhoods( $R, \epsilon$ )

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**Input:** Collection of sets  $R$ , distance threshold  $\epsilon$   
**Result:** Materialized neighborhoods of  $R$  w.r.t.  $\epsilon$

- 1  $I \leftarrow \text{Create-Index}(R, \epsilon)$ ;
- 2  $\text{pairs} \leftarrow \emptyset$  // Result set of similar pairs
- 3 **foreach**  $r \in R$  *in processing order* **do**
- 4      $M \leftarrow \text{Probe}(r, I, \epsilon)$  // candidates with prefix overlaps
- 5     **foreach**  $(s, po) \in M$  **do** //  $po \dots$  prefix overlap
- 6         **if**  $\text{Verify-Pair}(r, s, \epsilon, po)$  **then**
- 7              $\text{pairs} \leftarrow \text{pairs} \cup \{(r, s)\}$
- 8  $\text{neighborhoods} \leftarrow$  new associative array of size  $|R|$ ;
- 9 **foreach**  $(r, s) \in \text{pairs}$  **do**
- 10      $\text{neighborhoods}[r] \leftarrow \text{neighborhoods}[r] \cup \{s\}$ ;
- 11      $\text{neighborhoods}[s] \leftarrow \text{neighborhoods}[s] \cup \{r\}$ ;
- 12 **return**  $\text{neighborhoods}$

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**Algorithm 2:** Create-Index( $R, \epsilon$ )

---

**Input:** Collection of sets  $R$ , distance threshold  $\epsilon$   
**Result:** Inverted index of  $R$  w.r.t.  $\epsilon$

- 1  $I \leftarrow \emptyset$  // inv. index of set prefixes,  $I_{r[p]} \dots$  list of token  $r[p]$
- 2 **foreach**  $r \in R$  **do**
- 3      $\pi \leftarrow \lfloor \frac{\epsilon+2}{2} \rfloor$  // indexing prefix length of  $r$
- 4     **for**  $p \leftarrow 1$  **to**  $\pi$  **do**  $I_{r[p]} \leftarrow I_{r[p]} \cup \{r\}$ ;
- 5 **return**  $I$

---

### 3.2 Join-Clust: Materialized Neighborhoods

Join-Clust executes a set similarity self-join on  $R$  and materializes the  $\epsilon$ -neighborhoods in main memory. The self-join traverses all sets of  $r \in R$  in processing order and computes their lookahead neighbors,  $N_\epsilon^>(r)$ . The lookahead neighbors of  $r$  are appended to the list of  $r$ 's neighbors, and  $r$  is appended to the neighborhood lists of all  $s \in N_\epsilon^>(r)$ . After processing all sets, the neighborhood list of each set  $r \in R$  is complete and stores  $N_\epsilon(r)$ .

Next, standard DBSCAN (cf. Section 2.4) is executed to form clusters using the materialized neighborhoods. Algorithms 1–4 implement the similarity join with neighborhood materialization, index creation, probing, and efficient verification [29].

Mann et al. [29] found that the prefix-based index in combination with the length filter can be considered state of the art given an efficient verification procedure (which we use).

Figure 6 shows the asymmetric prefix index for our running example,  $\epsilon = 3$ ,  $\pi^i = 2$ . We probe  $r_8 = \{3, 4, 7, 8\}$  and look up the lists of the tokens 3, 4, 7, and 8 (the length of the probing prefix is  $\pi = 4$ ). Since we are only interested in the lookahead neighbors, i.e., all neighbors that follow  $r_8$  in processing order, we need to inspect the lists only starting from the point where  $r_8$  or a set  $r_i > r_8$  appears. The length filter does not prune any candidate in this example, and the candidate set is  $\{r_9\}$ . Since  $H(r_8, r_9) > \epsilon$ , the lookahead neighborhood of  $r_8$  is empty,  $N_\epsilon^>(r_8) = \emptyset$ .

In the context of the self-join,  $r_8$  will be retrieved as a lookahead neighbor of  $r_6$ , which is processed before  $r_8$ . Therefore, the neighborhood list of  $r_6$  will store  $r_8$  and vice versa.

Join-Clust produces fewer candidates than Sym-Clust and is therefore faster. However, the efficiency of Join-Clust comes at the cost of a larger memory footprint since all neighborhoods must be materialized.

---

**Algorithm 3:** Probe( $r, I, \epsilon$ )

---

**Input:** Probing set  $r$ , inv. index  $I$ , distance threshold  $\epsilon$   
**Result:** Set of candidates for  $r$  w.r.t.  $\epsilon$

//  $M$  maps a candidate  $s$  to its prefix overlap with  $r$

- 1  $M \leftarrow$  new associative array // candidates
- 2  $\pi \leftarrow \epsilon + 1$  // probing prefix length of  $r$
- 3  $lb_r \leftarrow |r| - \epsilon$  // size lower bound wrt.  $r$
- 4 **for**  $p \leftarrow 1$  **to**  $\pi$  **do**
- 5     **foreach**  $s \in I_{r[p]}$  *in proc. order* **do** // list of token  $r[p]$
- 6         **if**  $|s| < lb_r$  **then break** ;
- 7         **else** // add candidate
- 8             **if**  $s \notin M$  **then**  $M[s] \leftarrow 0$ ; // init.
- 9              $M[s] \leftarrow M[s] + 1$  // incr. overlap of  $(r, s)$
- 10 **return**  $M$

---

---

**Algorithm 4:** Verify-Pair( $r, s, \epsilon, po$ )

---

**Input:** Probing set  $r$ , candidate set  $s$ , distance threshold  $\epsilon$ , prefix overlap  $po$   
**Result:** True iff  $r$  and  $s$  are similar w.r.t.  $\epsilon$ , false otherwise

// cf. Mann et al. [29] for prefixes, equiv. overlaps, and Verify proc.

- 1  $\pi_r, \pi_s \leftarrow$  probing resp. indexing prefix length of  $r$  resp.  $s$ ;
- 2  $w_r, w_s \leftarrow$   $\pi_r$ - resp.  $\pi_s$ -th token in  $r$  resp.  $s$ ;
- 3  $t \leftarrow$  equivalent overlap for  $r, s$ , and  $\epsilon$ ;
- 4 **if**  $w_r < w_s$  **then**
- 5     **return**  $\text{Verify}(r, s, t, po, \pi_r + 1, po + 1)$
- 6 **return**  $\text{Verify}(r, s, t, po, po + 1, \pi_s + 1)$

---

*Complexity Analysis.* A neighborhood query is a constant-time lookup and a traversal of  $O(|N_\epsilon(r)|)$  neighbors. In the worst case, the join reports  $O(n^2)$  pairs. Consequently, materializing the neighborhoods takes  $O(n^2)$  time and space for  $n = |R|$ . The asymmetric prefix index requires only  $O(n)$  space and does not dominate memory usage.

## 4 THE SPREAD ALGORITHM

We present *Spread*, a novel time- and space-efficient solution for the density-based clustering problem. Spread leverages the effective asymmetric index and clusters all sets by traversing the sets in processing order. We identify key challenges that must be solved, discuss the algorithm, prove its correctness, analyze time and space complexity, and sketch a multi-core extension.

### 4.1 Key Challenges

Since Spread uses an asymmetric neighborhood index, a processing order,  $>$ , must be imposed on the data points, and an index lookup of query point  $r$  retrieves only the lookahead neighbors,  $N_\epsilon^>(r)$ . To achieve a correct clustering without materializing the neighborhoods, three key challenges must be solved.

In the following discussion, we assume that all sets of  $R$  are processed in processing order. When the *current set*  $r_i$  is to be processed, we know the core status of all preceding sets,  $r_j < r_i$ , but we do not know the core status of any unprocessed sets,  $r_k > r_i$ . We further assume that all sets  $r \in R$  that are directly density-reachable from any  $r_j$  that precedes  $r_i$  (i.e., are neighbors of a core point  $r_j < r_i$ ) are assigned to the same cluster as the core point  $r_j$ ; this may also include sets  $r_k > r_i$  that have not been processed yet.

*Core Status.* A set  $r_i$  is core if  $|N_\epsilon(r_i)| \geq \text{minPts}$ . Sym-Clust and Join-Clust have access to the full neighborhood,  $N_\epsilon(r_i)$ , thus deciding the core status of  $r_i$  is trivial. In contrast, Spread sees only the lookahead neighbors,  $N_\epsilon^\succ(r_i)$ . To identify the core status of  $r_i$ , however, additional knowledge about the size of the preceding neighborhood,  $N_\epsilon^\prec(r_i)$ , is required.

Consider probing  $r_i = r_5$  in our running example. According to our assumptions, the core status of sets  $r_1-r_4$  is known (only  $r_3$  is core), and all neighbors of  $r_3$  are in cluster  $C_3 = \{r_1, r_3, r_4, r_6, r_7\}$ . An index lookup of  $r_5$  returns  $N_\epsilon^\succ(r_5) = \{r_9, r_{10}\}$ . Since  $|N_\epsilon^\succ(r_5)| + 1 = 3 < 4 = \text{minPts}$  we cannot decide if  $r_5$  is core. In fact,  $r_5$  should be classified core since the full neighborhood is  $N_\epsilon(r_5) = \{r_2, r_5, r_9, r_{10}\}$  (cf. red circle in Figure 3).

*Border vs. Noise.* Assume that the current set  $r_i$  is a non-core point that is not assigned to any cluster. We need to decide if  $r_i$  is border or noise. A border point has at least one core point in its neighborhood. None of the preceding neighbors,  $r_j \in N_\epsilon^\prec(r_i)$ , is core, otherwise  $r_i$  would be assigned to the cluster of  $r_j$ . Thus,  $r_i$  is core iff one of the lookahead neighbors is core. Unfortunately, we do not know the core status of the lookahead neighbors and can therefore not label  $r_i$  as border or noise.

Assume a core point,  $r_k \in N_\epsilon^\succ(r_i)$ , among the lookahead neighbors of  $r_i$ . When  $r_k$  is processed,  $r_k$  will not see  $r_i$  in its lookahead neighborhood since  $r_i < r_k$ . Therefore,  $r_i$  will not be included into the cluster of  $r_k$  and will wrongly be classified noise. The challenge is to correctly decide the border status of  $r_i$  despite seeing only the lookahead neighbors of  $r_i$  and  $r_k$ .

In our running example,  $r_1$  is processed first. Thus, no core points are known and no clusters exist.  $N_\epsilon^\succ(r_1) = \{r_3\}$  and  $r_1$  remains noise (cf. blue circle in Figure 3). When the neighbor  $r_3$  of  $r_1$  is processed,  $r_3$  will be detected as a core point and start a new cluster. However, since  $r_3$  only sees its lookahead neighbors,  $N_\epsilon^\succ(r_3) = \{r_4, r_6, r_7\}$ ,  $r_1$  is not included into the cluster and is not detected as a border point.

*Disconnected Clusters.* Assume that the current set  $r_i$  is core and there is a core point  $r_j < r_i$  in a cluster  $C_j$ ,  $r_i \notin C_j$ . The current set  $r_i$  will assign all its lookahead neighbors to its cluster,  $C_i = C_i \cup N_\epsilon^\succ(r_i)$  ( $C_i$  can be a new cluster started by  $r_i$  or an existing cluster to which  $r_i$  belongs). Unfortunately, we cannot assume that  $C_i$  and  $C_j$  are indeed distinct clusters: there can be a core point  $r_k > r_i$  that density-reaches both  $r_i$  and  $r_j$ , i.e.,  $C_i$  and  $C_j$  should form a single cluster. In general, multiple subclusters of the same DBSCAN cluster may grow independently. The challenge is to identify subclusters that should be merged and to merge them efficiently.

We process the current set  $r_i = r_6$  in our running example. According to our assumptions, we know that  $r_3$  and  $r_5$  are core and we are aware of two clusters,  $C_3 = \{r_1, r_3, r_4, r_6, r_7\}$ ,  $C_5 = \{r_2, r_5, r_9, r_{10}\}$ . In addition, assume that we know that  $r_6$  is core. Then,  $r_6$  extends its current cluster,  $C_3$ , with its lookahead neighbors  $N_\epsilon^\succ(r_6) = \{r_8, r_{10}\}$ . Note that  $r_{10}$  is already part of cluster  $C_5$ . Since we do not know the core status of  $r_{10}$ , we cannot decide if  $C_5$  and  $C_3$  should be merged into a single cluster. If  $r_{10}$  is core,  $r_5$  and  $r_3$  are density-reachable from  $r_{10}$  and should be in the same cluster. If  $r_{10}$  is a border point, however, the clusters must not be merged, and  $r_{10}$  can be assigned to either  $C_5$  or  $C_3$ .

## 4.2 Data Structures

*Disjoint-Set.* The disjoint-set (or union-find) data structure maintains a dynamic collection of non-overlapping sets for  $n$

objects in  $O(n)$  space [10, 41]. A typical use case is the efficient computation of (minimum) spanning trees. It supports three operations: (1) For a given element  $u$ ,  $\text{make\_set}(u)$  creates a new (singleton) set that contains  $u$ . (2) The union( $u, v$ ) operation merges the two sets that contain  $u$  resp.  $v$  into a new set. (3)  $\text{find\_set}(u)$  returns the representative for the set that contains  $u$  or  $\infty$  if  $u$  is not found. The amortized worst-case time complexity is  $\Theta(\alpha(n))$  for all operations,  $\alpha(\cdot)$  being the inverse Ackermann function. In practice,  $\alpha(n)$  is considered a constant. In our setting, set elements are subclusters, and the disjoint-set data structure links subclusters that belong to the same DBSCAN cluster.

*Backlinks.* The backlinks data structure of a set  $r \in R$  is a collection of unique references to other sets  $s$  that precede  $r$ ,  $s < r$ . The backlinks  $bl$  support the add operation,  $bl \cup \{s\}$ , which adds a reference to a new set  $s$  in time  $O(1)$  (on average). Depending on the type of sets that are referenced in the backlinks, we distinguish core and non-core backlinks, denoted  $c\_bl$  and  $nc\_bl$ , respectively. We implement backlinks as unordered sets of integer identifiers.

## 4.3 The Algorithm

Algorithm 5 shows the pseudocode of Spread. We use the following notation:  $r$  is the current probing set,  $s > r$  is a lookahead neighbor, and  $x < r$  is a preceding neighbor. Initially, all sets are noise, i.e., their cluster identifier is  $-\infty$ ,  $\forall r \in R : r.cid = -\infty$ . Although we initialize all sets in Algorithm 5 explicitly (lines 3–4), this can also be done during indexing (cf. Algorithm 2).

*Algorithm Outline.* Spread proceeds in three main steps: (1) A counter and the processing order guarantee that the cardinality of the  $\epsilon$ -neighborhood is known when a set is processed despite using the asymmetric prefix index. (2) Each set is assigned to a subcluster solely based on its lookahead neighborhood. Subclusters of the same DBSCAN cluster are linked in a subcluster graph. Backlinks ensure that we do not miss border sets or links between subclusters. (3) Each connected component in the subcluster graph represents a DBSCAN cluster.

*Core Status.* A set  $r$  is core if  $|N_\epsilon(r)| \geq \text{minPts}$ . In Spread, however, only  $N_\epsilon^\succ(r)$  is computed. To capture the cardinality of  $N_\epsilon^\prec(r)$ , we store a density counter with each set  $r$ , denoted  $r.dens$ . Initially,  $\forall r \in R : r.dens = 1$ . For every lookahead neighbor  $s \in N_\epsilon^\succ(r)$ ,  $r.dens$  and  $s.dens$  are incremented (due to the symmetry of the distance). Core set identification is highlighted in green  $\square$ .

*Border vs. Noise.* A probing set  $r$  that is not core is a border set iff  $\exists y \in N_\epsilon(r) : y$  is core. Due to our processing order and the fact that only  $N_\epsilon^\succ(r)$  is computed, the existence of a core neighbor  $y$  may be unknown when  $r$  is probed. However, for each  $s \in N_\epsilon^\succ(r)$ , we know that  $r$  is part of  $N_\epsilon^\prec(s)$ . We store this information by adding  $r$  to the non-core backlinks  $nc\_bl[s]$  of each  $s \in N_\epsilon^\succ(r)$  (lines 31–33). Then, the first  $s \in N_\epsilon^\succ(r)$  that becomes core claims  $r$  (and all other unassigned sets in  $nc\_bl[s]$ ) as border point for its subcluster. If none of the neighbors  $s \in N_\epsilon^\succ(r)$  becomes core, then  $r$  remains noise. Lines 26–30 deal with a special case: If any  $s \in N_\epsilon^\succ(r)$  is already core when  $r$  is probed, then  $s$  claims  $r$  immediately without adding  $r$  to its non-core backlinks. The relevant code lines are marked in red  $\square$ .

*Subcluster Linkage.* If the probing set  $r$  is core and a core neighbor  $y$  is part of another subcluster, the subclusters of  $r$  and  $y$  must be linked in our subcluster graph. The subcluster graph represents all connected components of subclusters, each of which is

a DBSCAN cluster. We use the disjoint-set data structure  $ds$  to track the connected components. Two subclusters  $u, v$  are linked by  $ds.union(u, v)$ . We may not be able to determine if there is a set  $s \in N_\epsilon^\geq(r)$  that is core before  $s$  is probed. We use the core backlinks,  $c\_bl$ , to book-keep potential subclusters for linkage:  $r$  adds its subcluster identifier to  $c\_bl[s]$  of each  $s \in N_\epsilon^\geq(r)$  (lines 22-23). After  $N_\epsilon^\geq(r)$  has been processed, a link between the subcluster of  $r$  and every entry in  $c\_bl[r]$  is created (line 24). The special case when  $s$  is already core allows us to create the link immediately without using core backlinks (lines 20-21). Linkage is only required if two subclusters coalesce (condition in line 19). Otherwise,  $r$  simply claims  $s \in N_\epsilon^\geq(r)$  for its subcluster (lines 17-18). Linkage of subclusters is highlighted in blue  $\square$ .

All backlinks of  $r$  are released after  $r$  has been processed to save memory (line 34). The subcluster graph in  $ds$  is used to assign consistent cluster IDs in a final scan over  $R$  (lines 35-36).

#### 4.4 Correctness

We show that Algorithm 5 partitions  $R$  into DBSCAN clusters (cf. Definition 2.1). Set  $r_i \in R$  is the  $i$ -th set of  $R$  in processing order. We prove the correctness by induction over  $i$  and increasing subsets  $R^i \subseteq R$ .  $R^0 = \emptyset$ ,  $R^i = R^{i-1} \cup \{r_i\} \cup N_\epsilon^\geq(r_i)$  for  $1 \leq i \leq n = |R|$ , thus  $R^n = R$ . Due to space constraints, we omit the full proofs and only provide the invariants that must be shown.

*Core Status.* The core status of set  $r_i$  is determined in the  $i$ -th iteration of the main loop.  $r_i$  is core if  $\minPts \leq |N_\epsilon(r_i)| = 1 + |N_\epsilon^\leq(r_i)| + |N_\epsilon^\geq(r_i)|$ . In line 5,  $r_i.dens = 1 + |N_\epsilon^\leq(r_i)|$ . Lines 6-11 compute  $N_\epsilon^\geq(r_i)$ . The index lookup in line 6 returns candidate set  $M$ ,  $N_\epsilon^\geq(r_i) \subseteq M \subseteq \{s \mid s > r_i\}$ . Every set  $s \in M$  is verified in line 9 such that  $N_\epsilon^\geq(r_i)$  is available starting from line 12.

LEMMA 4.1. *Algorithm 5 correctly identifies all core sets in  $R$ .*

PROOF SKETCH. We show that at the start of the  $i$ -th iteration in line 5, for all  $r_k$  and  $r_j$ ,  $1 \leq k < i \leq j$  the following invariants hold: (I1)  $r_k.dens = |N_\epsilon(r_k)|$ ; (I2)  $r_j.dens = 1 + |\{r_k \mid r_j \in N_\epsilon^\geq(r_k)\}|$ , i.e.,  $r_i.dens = 1 + |N_\epsilon^\leq(r_i)|$ . Further, (I3) in line 12 of the  $i$ -th iteration,  $r_i.dens = |N_\epsilon(r_i)|$ , i.e., Algorithm 5 correctly identifies the core status of  $r_i$ .  $\square$

*Border vs. Noise.* Lines 25-33 cover the case that  $r_i$  is not core. If any  $s \in N_\epsilon^\geq(r_i)$  qualifies as core,  $s$  claims  $r_i$ . Otherwise,  $r_i$  is stored in the non-core backlinks  $nc\_bl[s]$  of every  $s \in N_\epsilon^\geq(r_i)$  (lines 31-33). The next core neighbor in processing order claims  $r_i$  (lines 14-15) such that all border sets are assigned to a cluster.

LEMMA 4.2. *Algorithm 5 correctly clusters all border sets in  $R$ .*

PROOF SKETCH. At the start of the  $i$ -th iteration, the following invariant holds for all border sets  $r_k \in \mathcal{B}$ ,  $1 \leq k < i$ : if  $r_k$  is not stored in  $nc\_bl[s]$  for any  $s \in N_\epsilon^\geq(r_k)$ ,  $s = r_i$  or  $s > r_i$ , then  $r_k$  is assigned to the cluster of a core point in its neighborhood.  $\square$

*Subcluster Linkage.* Lines 12-24 cover the case that  $r_i$  is core. Each core point may form a subcluster on its own or together with other core points. We must ensure that all subclusters of the same DBSCAN cluster are linked in the disjoint-set,  $ds$ .

LEMMA 4.3. *Algorithm 5 correctly links all subclusters in  $R$ .*

PROOF SKETCH. At the start of the  $i$ -th iteration, the following invariant holds for all core neighbors  $c \in CN(r_k) = N_\epsilon(r_k) \cap C$  of a core set  $r_k \in C$ ,  $1 \leq k < i$ : (a)  $c$  and  $r_k$  have the same cluster representative (in  $ds$ ), or (b)  $c$  is stored in some  $c\_bl[s]$ ,  $s \in N_\epsilon^\geq(r_k)$ ,  $s = r_i$  or  $s > r_i$ .  $\square$

#### Algorithm 5: Spread( $R, \epsilon, \minPts$ )

---

**Input:** Collection of sets  $R$ , distance threshold  $\epsilon$ ,  
min. density  $\minPts$

**Result:** A correct DBSCAN clustering of  $R$  w.r.t.  $\epsilon, \minPts$

```

1  $ds \leftarrow$  new disjoint-set;  $nc\_bl, c\_bl \leftarrow$  new backlinks;
2  $I \leftarrow$  Create-Index( $R, \epsilon$ );
3 foreach  $r \in R$  do
4    $r.dens \leftarrow 1$ ;  $r.cid \leftarrow -\infty$ ;  $ds.make\_set(r.id)$ ;
5 foreach  $r \in R$  in processing order do
6    $M \leftarrow$  Probe( $r, I, \epsilon$ );
7    $N_\epsilon^\geq(r) \leftarrow \emptyset$ ;
8   foreach  $(s, po) \in M$  do //  $po \dots$  prefix overlap
9     if Verify-Pair( $r, s, \epsilon, po$ ) then
10        $r.dens \leftarrow r.dens + 1$ ;  $s.dens \leftarrow s.dens + 1$ ;
11        $N_\epsilon^\geq(r) \leftarrow N_\epsilon^\geq(r) \cup \{s\}$ ;
12   if  $r.dens \geq \minPts$  then //  $r$  is core
13     if  $r.cid = -\infty$  then  $r.cid \leftarrow r.id$ ;
14     foreach  $x \in nc\_bl[r]$  do // claim border sets  $x < r$ 
15       if  $x.cid = -\infty$  then  $x.cid \leftarrow r.cid$ ;
16     foreach  $s \in N_\epsilon^\geq(r)$  do //  $s > r$ 
17       if  $s.cid = -\infty$  then // claim unclaimed  $s > r$ 
18          $s.cid \leftarrow r.cid$ 
19       else if  $r.cid \neq s.cid$  then //  $s$  already claimed
20         if  $s.dens \geq \minPts$  then //  $s$  is core
21            $ds.union(r.cid, s.cid)$  // link subclusters
22         else // remember core neighbor  $r$ 
23            $c\_bl[s] \leftarrow c\_bl[s] \cup \{r.cid\}$ 
24     foreach  $x \in c\_bl[r]$  do  $ds.union(r.cid, x)$ ;
25   else //  $r$  is not core, i.e.,  $r.dens < \minPts$ 
26     if  $r.cid = -\infty$  then // claim potential border set  $r$ 
27       foreach  $s \in N_\epsilon^\geq(r)$  do
28         if  $s.dens \geq \minPts$  then //  $s$  is core
29           if  $s.cid = -\infty$  then  $s.cid \leftarrow s.id$ ;
30            $r.cid \leftarrow s.cid$ ; break;
31     if  $r.cid = -\infty$  then // remember potential border set  $r$ 
32       foreach  $s \in N_\epsilon^\geq(r)$  do
33          $nc\_bl[s] \leftarrow nc\_bl[s] \cup \{r\}$ 
34   release  $c\_bl[r]$  and  $nc\_bl[r]$  // not needed anymore
35 foreach  $r \in R$  do // final assignment of cluster IDs
36   if  $r.cid \neq -\infty$  then  $r.cid \leftarrow ds.find\_set(r.cid)$ ;

```

---

THEOREM 4.4. *Algorithm 5 returns a correct set clustering  $\Gamma = \{C_1, C_2, \dots, C_k\}$  of  $R$  according to Definition 2.1.*

PROOF SKETCH. By Lemmata 4.1-4.3 and due to our final scan over  $R$  (lines 35-36),  $x.cid = ds.find\_set(x.cid)$  holds for all  $x \in R$ . Initially,  $x.cid = -\infty$  for all  $x \in R$ . The cluster IDs are updated only for border and core sets. Consequently,  $x.cid = -\infty$  holds for all  $x \in R \setminus (C \cup \mathcal{B}) \equiv \mathcal{N}$ , i.e., also noise is correctly identified.  $\square$

## 4.5 Complexity Analysis

*Memory.* The asymmetric prefix index requires  $O(n)$  space. In addition, Spread maintains the following data structures. (i) A density counter for each set  $r \in R$  requires  $O(n)$  space. (ii) A disjoint-set data structure with at most  $O(n)$  entries, i.e., the disjoint-set structure requires  $O(n)$  space [41]. (iii) In the worst case, we allocate two backlink structures for each  $r \in R$ , i.e.,  $O(n)$  backlinks. We release  $c\_bl[r]$  and  $nc\_bl[r]$  after probing  $r$ . Backlinks are only extended in lines 23 and 33. However, both lines are executed iff  $\exists s \in N_\epsilon^>(r) : s$  is core. Set  $s$  is core iff  $s.dens \geq \text{minPts}$ , and the density is updated for every neighbor, therefore any backlink holds at most  $\text{minPts}$  entries. As a result, no more than  $O(n \cdot \text{minPts})$  entries are allocated, thus requiring  $O(n)$  space since  $\text{minPts}$  and  $\epsilon$  are constants. *Runtime.* For each  $r \in R$ , we process  $O(|N_\epsilon^>(r)|)$  neighbors and the backlinks of  $r$  if it is core. Recall that the disjoint-set operations take constant time. Therefore, the final for-loop (lines 35–36) runs in  $O(n)$  time. Overall, Spread runs in  $O(n^2)$  time and  $O(n)$  space.

## 4.6 Multi-core Extension

Spread is designed as a single-core algorithm. We sketch an extension to multi-core processors that requires little synchronization between threads. Our extension is based on the observation that Spread spends most of the runtime in neighborhood computations (lines 6–11). While for some datasets the neighborhood computation accounts for only about half of the overall runtime (e.g., 55% for ORKUT,  $\epsilon = 3$ ), for the configuration with the highest runtime in our experiments (CELONIS1,  $\epsilon = 5$ ), Spread spends over 99% of the runtime in computing the neighborhoods.

We distribute the workload to  $k + 1$  threads,  $T_1, T_2, \dots, T_{k+1}$ . Threads  $T_1 - T_k$  are responsible for the neighborhood computations (lines 6–11),  $T_{k+1}$  performs the actual clustering (lines 12–34). The runtime of the other steps in the algorithm is negligible.

*Neighborhood Computation.* Let  $r_i \in R$ ,  $1 \leq i \leq |R|$  be the  $i$ -th set of  $R$  in processing order. Thread  $T_j$ ,  $1 \leq j \leq k$ , computes the neighborhoods  $N_\epsilon^>(r_i)$  of all  $r_i$  with  $j = i \bmod k$  (i.e., round robin). Each thread processes the assigned sets  $r_i$  in processing order (i.e., increasing values of  $i$ ). The neighborhood computation in Algorithm 5 is interleaved with updating the density counters of  $r_i$  and its neighbors. Only this step requires synchronization (e.g., using atomic writes) since multiple threads may access the same counter concurrently. We do not expect congestions since the density updates are distributed over all neighbors.

*Cluster Scan.* Thread  $T_{k+1}$  scans the sets in processing order and performs the steps in lines 12–34 (maintain backlinks and disjoint-set, assign preliminary cluster IDs). After processing a set  $r_i$ , the memory for the neighbors of  $r_i$  is released.

*Synchronization.* We need to make sure that  $T_{k+1}$  processes set  $r_i$  only after  $r_i$ 's neighbors have been computed. This can be achieved with a lock (implemented as condition variable<sup>2</sup>) on  $r_i$  that is held by  $T_j$ ,  $j \leq k$ , until the neighborhood of  $r_i$  is computed.  $T_{k+1}$  needs to get the lock on  $r_i$  before processing it.

*Memory.*  $T_{k+1}$  releases the neighbors after processing them. If the parallel neighborhood computation is faster than  $T_{k+1}$ , the precomputed neighborhoods will fill up the memory. This is avoided with a shared counter that is incremented by  $T_1 - T_k$  (when they process a new set  $r_i$ ) and is decremented by  $T_{k+1}$  (after processing  $r_i$ ). The neighborhood computation of  $r_i$  is postponed until the counter is below some threshold that bounds the number of concurrently materialized lookahead neighborhoods.

<sup>2</sup>A queue of threads waiting for a condition to become true.

Table 2: Characteristics of datasets.

Dataset	Coll. Size	Set Size		Univ. Size
		avg.	max.	
BMS-POS <sup>4</sup>	$3.2 \cdot 10^5$	9.3	164	$1.7 \cdot 10^3$
FLICKR <sup>5</sup>	$1.2 \cdot 10^6$	10.1	102	$8.1 \cdot 10^5$
KOSARAK <sup>6</sup>	$6.1 \cdot 10^5$	11.9	$2.5 \cdot 10^3$	$4.1 \cdot 10^4$
LIVEJ <sup>7</sup>	$3.1 \cdot 10^6$	36.4	300	$7.5 \cdot 10^6$
ORKUT <sup>7</sup>	$2.7 \cdot 10^6$	119.7	$4.0 \cdot 10^4$	$8.7 \cdot 10^6$
SPOT <sup>8</sup>	$4.4 \cdot 10^5$	12.8	$1.2 \cdot 10^4$	$7.6 \cdot 10^5$
CELONIS1	$8.2 \cdot 10^6$	20.3	91	$1.2 \cdot 10^4$
CELONIS2	$2.6 \cdot 10^6$	22.1	130	$3.5 \cdot 10^3$

## 5 EXPERIMENTAL EVALUATION

*Algorithms.* We compare our solution, Spread, against the two baseline approaches Sym-Clust and Join-Clust (cf. Section 3). All algorithms are single-threaded C++ implementations (2017 standard). Our implementations of Spread, Join-Clust, and the index of Sym-Clust follow the guidelines by Mann et al. [29], e.g., regarding symmetric and asymmetric prefix index, candidate generation, and optimized prefix-based verification.

*Datasets.* We execute all experiments on 13 real-world datasets: (a) Nine of the datasets where previously used for benchmarking set similarity joins [16, 29]: BMS-POS, DBLP, ENRON, FLICKR, KOSARAK, LIVEJ, NETFLIX, ORKUT, and SPOT. For a description of the datasets and preprocessing instructions<sup>3</sup> we refer to Mann et al. [29]. (b) Four large real-world datasets from the process mining domain, CELONIS1–4, that store one set per process. Compared to most datasets of the join benchmark, the universe size of these datasets is rather small. Table 2 summarizes important characteristics of our benchmark data.

Due to space constraints we omit detailed results for the following datasets: (a) DBLP, ENRON, and NETFLIX show very low runtimes ( $< 4s$ ) and a small and stable memory footprint ( $< 1\text{GiB}$ ) for all algorithms and configurations. (b) CELONIS3–4 show results similar to the other process mining datasets.

*Parameters.* The algorithms take two parameters: the neighborhood radius,  $\epsilon$ , and the density,  $\text{minPts}$ . Typically, density-based clustering is sensitive to  $\epsilon$  and quite robust to  $\text{minPts}$ . In our experiments, we vary both parameters:  $\epsilon \in \{2, 3, 4, 5\}$  and  $\text{minPts} \in \{2, 4, 8, 16, 32, 64, 128\}$  (defaults in bold font).

*Environment.* All experiments have been conducted on a 64-bit machine with 2 physical Intel Xeon E5-2630 v3 CPUs, 2.40GHz, 8 cores each (i.e., 16 logical processors, hyper-threading disabled). The cores share a 20MiB L3 cache and have another 256KiB of independent L2 cache. The system has 96GiB of RAM and runs Debian 10 Buster (Linux 4.19.0-12-amd64 #1 SMP Debian 4.19.152-1 (2020-10-18)). Our code is compiled with clang<sup>9</sup> version 7, highest optimization level ( $-O3$ ). The runtime is measured with `clock_gettime`<sup>10</sup> at process level, memory usage is the heap

<sup>3</sup><http://ssjoin.dbresearch.uni-salzburg.at/datasets.html>

<sup>4</sup>BMS-POS: <http://www.kdd.org/kdd-cup/view/kdd-cup-2000> [52]

<sup>5</sup>FLICKR: Bouros et al. [6]

<sup>6</sup>KOSARAK: <http://fimi.uantwerpen.be/data/>

<sup>7</sup>LIVEJ, ORKUT: <http://socialnetworks.mpi-sws.org/data-imc2007.html> [30]

<sup>8</sup>SPOT: Pichl et al. [33]

<sup>9</sup><https://releases.lldvm.org/7.0.0/tools/clang/docs/ReleaseNotes.html>

<sup>10</sup>[https://man7.org/linux/man-pages/man2/clock\\_gettime.2.html](https://man7.org/linux/man-pages/man2/clock_gettime.2.html)

Table 3: Index & cluster statistics for  $\epsilon = 3$ ,  $\text{minPts} = 16$ .

(a) BMS-POS.			
	Candidates	True Positives	Clusters
Sym-Clust	$3.9 \cdot 10^9$	$38.0 \cdot 10^6$	1
Join-Clust	$640.0 \cdot 10^6$	$38.0 \cdot 10^6$	1
Spread	$640.0 \cdot 10^6$	$38.0 \cdot 10^6$	1
(b) KOSARAK.			
	Candidates	True Positives	Clusters
Sym-Clust	$40.7 \cdot 10^9$	$2.8 \cdot 10^9$	5
Join-Clust	$7.0 \cdot 10^9$	$2.8 \cdot 10^9$	5
Spread	$7.0 \cdot 10^9$	$2.8 \cdot 10^9$	5
(c) CELONIS1.			
	Candidates	True Positives	Clusters
Sym-Clust	$644.6 \cdot 10^9$	$7.4 \cdot 10^6$	5,075
Join-Clust	$131.5 \cdot 10^9$	$7.4 \cdot 10^6$	5,075
Spread	$131.5 \cdot 10^9$	$7.4 \cdot 10^6$	5,075

peak of Linux memusage<sup>11</sup> (using LD\_PRELOAD). A single instance is executed at a time with no other load on the machine.

## 5.1 Index & Cluster Statistics

We compare the number of candidates, true positives, and the number of clusters. The numbers are sums over all region queries. Table 3 shows the results obtained for BMS-POS, KOSARAK, and CELONIS1. We observe that Spread produces exactly the same number of candidates as Join-Clust since both solutions use the asymmetric index. Sym-Clust generates significantly more candidates due to the symmetric prefix index and the symmetric distance computations. For CELONIS1, Spread and Join-Clust verify about 5 times fewer candidates compared to Sym-Clust.

## 5.2 Runtime Efficiency

We measure the overall runtime, i.e., the CPU time that is required to cluster all sets into DBSCAN clusters (excluding the time to load the data from disk). Figure 7 shows the results for varying  $\epsilon$  ( $\text{minPts} = 16$ ). We observe that Sym-Clust is not competitive in terms of overall runtime in most cases. For all datasets, except KOSARAK and SPOT, the runtime of Sym-Clust increases much faster with  $\epsilon$  than observed for Join-Clust and Spread. This is mainly due to the use of the symmetric prefix index (more candidates) and redundant computations (symmetric pairs).

Our experiments reveal that Join-Clust suffers from the following issues: (i) High runtimes for LIVEJ, ORKUT, and SPOT due to the expensive neighborhood materialization. (ii) Join-Clust runs out of memory for many instances (missing points in plots), in particular for FLICKR (any  $\epsilon$ ), KOSARAK ( $\epsilon \geq 4$ ), LIVEJ, ORKUT, and SPOT ( $\epsilon \geq 3$ ).

Spread outperforms its competitors in most settings and is competitive with Join-Clust otherwise (cf. Figures 7a, 7g, and 7h). For CELONIS1 and CELONIS2, Spread outperforms Sym-Clust by almost an order of magnitude and is competitive with Join-Clust.

Figure 8 shows the runtime results for varying  $\text{minPts}$  values ( $\epsilon = 3$ ). We observe that the runtime of all three solutions is quite robust to  $\text{minPts}$ . The insights are similar for all datasets and values of  $\epsilon$ . We include the plots for BMS-POS and KOSARAK.

<sup>11</sup><https://man7.org/linux/man-pages/man1/memusage.1.html>

## 5.3 Memory Efficiency

We study the memory usage of Join-Clust, Sym-Clust, and Spread. All three solutions store (i) the collection, (ii) the inverted index, (iii) the candidates, and (iv) the result of a region query on the heap. The symmetric prefix index of Sym-Clust is larger than the asymmetric index, but still linear in the collection size. Sym-Clust generates more candidates than Join-Clust and Spread (cf. Section 5.1), which both use the asymmetric prefix index. Join-Clust materializes all neighborhoods in main memory. Sym-Clust and Spread materialize only a single neighborhood at a time. Spread stores also backlinks and the disjoint-set in main memory.

Figure 11 shows our results for varying  $\epsilon$  ( $\text{minPts} = 16$ , y-axis log scale). Join-Clust runs out of memory for many instances (cf. Section 5.2). The neighborhood materialization in Join-Clust can be memory intensive even for small values of  $\epsilon$ . We observe different growth rates with increasing radius  $\epsilon$ , which we attribute to the different neighborhood sizes. The memory consumption of Sym-Clust is significantly lower and robust to varying  $\epsilon$ . Spread shows a similar behavior. In some cases (e.g., LIVEJ, ORKUT), Spread occupies even less memory than Sym-Clust. When few backlinks are materialized, the smaller asymmetric prefix index of Spread outweighs the storage overhead for the backlinks.

Figure 12 shows the memory usage over  $\text{minPts}$  ( $\epsilon = 3$ , log-log scale). The memory consumption of Sym-Clust and Join-Clust is stable w.r.t. increasing values of  $\text{minPts}$ , while the memory usage of Spread slightly increases. This is due to the number of concurrently stored backlinks: the larger  $\text{minPts}$ , the higher the chance that a succeeding core neighbor is not yet classified, which triggers the creation of a backlink entry. The memory grows slowly with increasing  $\text{minPts}$  and does not limit the scalability of Spread. We include the results for BMS-POS and KOSARAK,  $\epsilon = 3$ ; other datasets and  $\epsilon$  values show similar results.

*Backlinks Peak.* We evaluate the effect of releasing the backlinks of a set in Spread after the set has been processed (cf. line 34, Algorithm 5). Figures 10 and 14 show the peak number of allocated backlinks relative to the maximum number of backlinks for varying  $\epsilon$  ( $\text{minPts} = 16$ ) and  $\text{minPts}$  ( $\epsilon = 3$ ), respectively. Since two backlink structures, core (green) and non-core (orange), are maintained for each set in  $R$ , at most  $2|R|$  backlinks can be allocated (light blue). Deallocating the backlinks of probed sets is highly effective: Only a small fraction of the maximum number of backlinks is allocated at any point in time. For increasing values of  $\epsilon$  and  $\text{minPts}$  also the number of allocated backlinks grows.

## 5.4 Scalability

We evaluate the scalability of Spread and its competitors to increasing data sizes. To this end, we increase the size of BMS-POS and KOSARAK using the procedure of Vernica et al. [42]. This approach does not affect the token universe, and the number of similar pairs in the dataset increases linearly with the data size.

Figure 9 (runtime) and Figure 13 (main memory) report the results for our default parameter setting. Spread shows runtimes similar to Join-Clust and outperforms Sym-Clust by a factor of about 12 (BMS-POS) resp. 5.7 (KOSARAK) on the largest dataset ( $\times 16$ ). As we increase BMS-POS by a factor of 16, the runtime increases by a factor of 195 for Spread, 204 for Join-Clust, and 460 for SymClust. The memory grows linearly for all measured data points and increases by a factor of about 2 when we double the data size. Join-Clust requires 18-25 (BMS-POS) resp. 499-569 (KOSARAK) times more memory than its competitors and runs out of memory on KOSARAK except for the  $\times 1$  dataset.

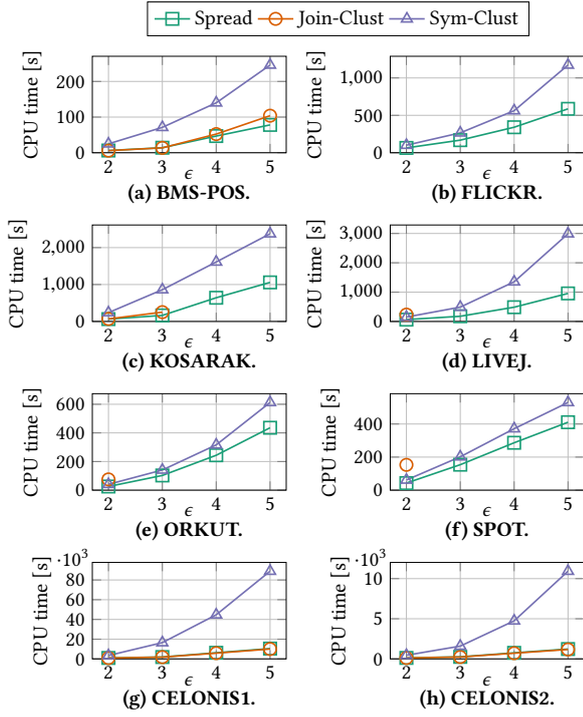


Figure 7: Runtime over  $\epsilon$ ,  $\text{minPts} = 16$ .

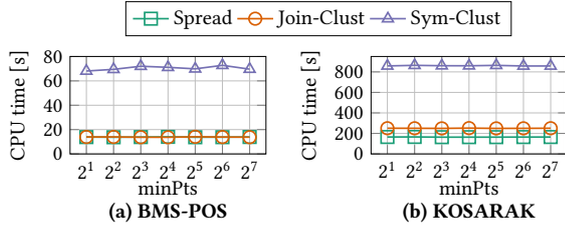


Figure 8: Runtime over  $\text{minPts}$ ,  $\epsilon = 3$ .

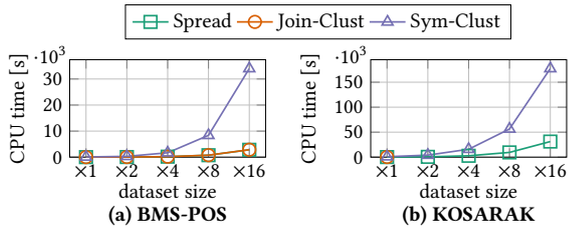


Figure 9: Runtime over data size,  $\epsilon = 3$ ,  $\text{minPts} = 16$ .

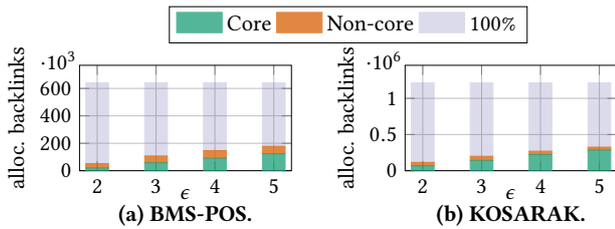


Figure 10: Backlinks peak over  $\epsilon$ ,  $\text{minPts} = 16$ .

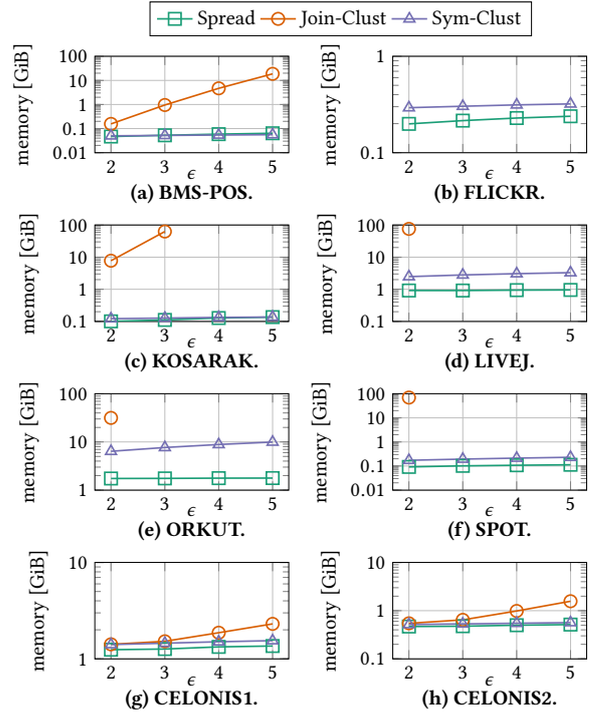


Figure 11: Main memory over  $\epsilon$ ,  $\text{minPts} = 16$ .

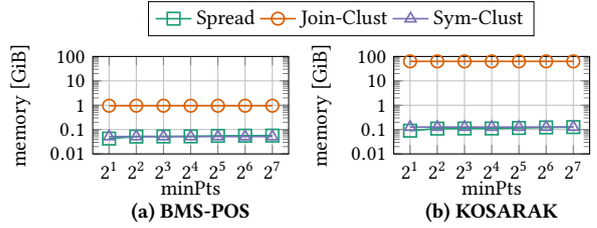


Figure 12: Main memory over  $\text{minPts}$ ,  $\epsilon = 3$ .

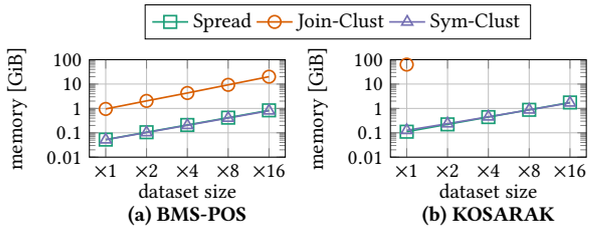


Figure 13: Main memory over data size,  $\epsilon = 3$ ,  $\text{minPts} = 16$ .

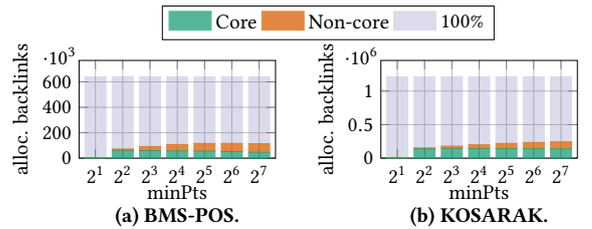


Figure 14: Backlinks peak over  $\text{minPts}$ ,  $\epsilon = 3$ .

Summarizing, Spread clearly outperforms Sym-Clust in runtime (by a factor of 5-12) and Join-Clust in memory usage (by more than an order of magnitude) as we increase the data size.

## 6 RELATED WORK

*Indexes for Sets.* Most set similarity joins operate on an inverted list index that maps signatures to candidate sets. Various signatures have been proposed [2, 8, 40, 45]. Prefixes [8] in conjunction with the length filter [1] have been shown to prune sets effectively. More sophisticated filters include positional and suffix filter [49], the removal filter [35], the position-enhanced length filter [28], and the adaptive prefix filter [44]. Wang et al. [46] leverage the similarity of the sets in an  $\epsilon$ -neighborhood to reduce the overall number of false positives. Dong et al. [13] propose a size-aware algorithm that runs in  $o(n^2) + O(k)$  time for  $k$  result pairs. Qin and Xiao [34] propose the pigeonring, a generalization of the pigeonhole principle that yields stronger constraints. Indexing and join techniques for sets have been studied extensively in the single-machine [29] and the distributed context [16].

Most of these approaches focus on self-joins, which order the sets and compute the lookahead neighborhood to avoid symmetric distance computations. In our work, we use the prefix filter, but any of the other asymmetric indexes is applicable.

*Efficient Region Queries.* Ester et al. [15] propose the first exact DBSCAN algorithm with  $O(n \log n)$  runtime for vectors of arbitrary dimension.  $O(n \log n)$  runtime holds for a small number of neighbors (compared to  $n$ ) and an index with  $O(\log n)$  lookup time. Henceforth, efficient region query computation has been of great interest and many improvements have been proposed. Brecheisen et al. [7] use minPts-nearest neighbor queries to identify core points and postpone the other distance computations until the distances are required to get a correct DBSCAN clustering. The proposed *Xseedlist* data structure is designed for expensive distance functions and assumes a cheap but selective filter. These assumptions do not hold for sets: The verification (i.e., distance computation) of candidate pairs has shown to be highly efficient [29] (a small number of integer comparisons). Brecheisen et al. must insert the candidates into the *Xseedlist* data structure, which maintains sorted lists of candidates. Due to the expensive sorting procedure, we do not expect *Xseedlist* to improve the DBSCAN algorithm for sets. TI-DBSCAN [25] exploits the triangle inequality to reduce the search space of region queries. The solution is not index-based, sorts the points w.r.t. a reference point, and shifts a window of size  $2\epsilon$  over the sorted points. The reference point is the point with minimal values in all dimensions. This is equivalent to the empty set, and our processing order in combination with the prefix index for sets subsumes this technique. Patwary et al. [32] introduce PARDICLE, a parallel approximate density-based clustering algorithm for Euclidean space. Its aim is to reduce the neighborhood computation time by sampling high-density regions. Kumar and Reddy [26] propose a new graph-based index structure called Groups. It discovers groups of patterns in two scans over the dataset and applies a standard DBSCAN afterwards. Groups accelerates region queries by pruning noise points effectively. This technique assumes Euclidean distance and does not consider Hamming distance or other set similarity measures. Recently, Jiang et al. [24] proposed SNG-DBSCAN, which prevents the computation of the full  $\epsilon$ -neighborhood graph via subsampling its edges. This results in  $O(sn^2)$ -time complexity with  $s$  being the sampling rate. Under certain distribution assumptions, SNG-DBSCAN has been shown

to preserve the exact  $\epsilon$ -neighborhood graph for  $s \approx (\log n)/n$  with  $O(n \log n)$  runtime.

*DBSCAN Techniques.* Yang et al. [51] propose the distributed DBSCAN-MS clustering algorithm for metric spaces. DBSCAN-MS uses pivots to map the data from metric space to vector space, where it is partitioned in order to be distributed. A local DBSCAN is then executed on each partition. Our solution does not rely on the metric properties of set distances, but uses specialized set indexes. However, our techniques may be leveraged in the context of DBSCAN-MS, where the data points are ordered by one of the dimensions for efficient neighborhood queries.

Patwary et al. [31] propose PDSDBSCAN, a parallel DBSCAN algorithm that uses the disjoint-set data structure to connect data points into clusters. We only insert links between subclusters into the disjoint-sets structure, while PDSDBSCAN inserts a link for each neighbor, rendering the number of required union operations a bottleneck for this approach.

Böhm et al. [3] use a block-nested loop join and buffer the join result to reduce the number of block accesses required to compute  $\epsilon$ -neighborhoods. CUDA-DClust [4] is a GPU-based solution that splits clusters into chains that are expanded from different starting points in parallel. In order to merge chains into clusters, a quadratic-size bit matrix is used. We maintain only a linear number of links and leverage disjoint-sets to merge clusters. Incremental DBSCAN algorithms [14] deal with updates on an existing clustering. Similar to our approach, these techniques may need to merge clusters when new points are inserted. None of the above solutions supports asymmetric neighborhood indexes.

Numerous parallel and distributed algorithms [9, 11, 18–21, 23, 36, 38, 47, 50] as well as approximations [17, 27, 43, 48] have been proposed. We present an exact, single-core solution for sets.

## 7 CONCLUSION

In this paper, we have investigated clustering techniques for large collections of sets. Our work was motivated by an application in process mining that models processes as sets to assess their similarity. We have shown that the solutions that are currently available, Sym-Clust and Join-Clust, are not satisfying: Sym-Clust is slow since it cannot use effective asymmetric set indexes, while Join-Clust is infeasible for many settings due to its excessive memory usage. We introduced a novel, density-based clustering algorithm, Spread, that can process data points in any user-defined order and is therefore fit for the use with asymmetric indexes. Spread combines the best of both worlds: It uses the effective asymmetric index of Join-Clust, but like Sym-Clust does not need to materialize the neighborhoods. We introduced so-called backlinks to guarantee a correct DBSCAN clustering and showed the correctness of our approach. To the best of our knowledge, Spread is the first DBSCAN-compliant algorithm that uses an asymmetric index and runs in linear space.

Spread uses the index as a black box and works with any data type. Interesting future work includes evaluating the performance of Spread for vector data, where candidates are generated using a sliding window that is shifted along one dimension. The data points in the window are candidates, i.e., the window simulates an asymmetric index for Spread.

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