# TI-DBSCAN: Clustering with DBSCAN by Means of the Triangle Inequality

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Abstract. Grouping data into meaningful clusters is an important data mining task. DBSCAN is recognized as a high quality density-based algorithm for clustering data. It enables both the determination of clusters of any shape and the identification of noise in data. The most time-consuming operation in DBSCAN is the calculation of a neighborhood for each data point. In order to speed up this operation in DBSCAN, the neighborhood calculation is expected to be supported by spatial access methods. DBSCAN, nevertheless, is not efficient in the case of high dimensional data. In this paper, we propose a new efficient TI-DBSCAN algorithm and its variant TI-DBSCAN-REF that apply the same clustering methodology as DBSCAN. Unlike DBSCAN, TI-DBSCAN and TI-DBSCAN-REF do not use spatial indices; instead they use the triangle inequality property to quickly reduce the neighborhood search space. The experimental results prove that the new algorithms are up to three orders of magnitude faster than DBSCAN, and efficiently cluster both low and high dimensional data.

# 1 Introduction

Grouping data into meaningful clusters is an important data mining task. The quality of clustering depends on a used algorithm. The DBSCAN algorithm (Density-Based Spatial Clustering of Applications with Noise) [3] is recognized as a high quality scalable algorithm for clustering low dimensional data. The most time-consuming operation in DBSCAN is the calculation of a neighborhood for each data point. In order to speed up this operation in DBSCAN, it is expected to be supported by spatial access methods such as R\*-tree [1] (R-tree [4]). DBSCAN, nevertheless, is not able to cluster high dimensional data efficiently. A method for improving the performance of DBSCAN based on early removal of core points has been offered in [6]. There, the carried out experiments showed that using the proposed method speeded up DBSCAN's performance by 50%.

In this paper, we propose a new efficient TI-DBSCAN algorithm and its variant TI-DBSCAN-REF that apply the same clustering methodology as DBSCAN. Unlike DBSCAN, TI-DBSCAN and TI-DBSCAN-REF do not use spatial indices; instead they use the triangle inequality property to quickly reduce the neighborhood search space. To the best of our knowledge, our proposal is the first one that relates to a density-based clustering; the other trials of using the triangle inequality in clustering

were related to k-means algorithm [2][7] and hierarchical algorithms following the results presented in [7].

The paper has the following layout. Section 2 recalls the notion of a cluster and noise according to [3]. In Section 3, we a offer theoretical basis of our approach to optimizing DBSCAN-like clustering. In Section 4, we propose the TI-DBSCAN algorithm and its modification TI-DBSCAN-REF. Section 5 reports the performance of TI-DBSCAN, TI-DBSCAN-REF as well as the performance of DBSCAN. Section 6 concludes the obtained results.

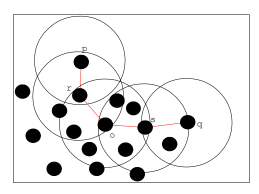
### 2 Basic Notions

In the context of the DBSCAN algorithm [3], a cluster is an area of high density. Data points in a low density area constitute noise. A point in space is considered a member of a cluster if there is a sufficient number of points within a given distance. In the sequel, the distance between two points p and q will be denoted by distance(p,q). Please, note that one may use a variety of distance metrics. Depending on an application, one metric may be more suitable than the other. In particular, if Euclidean distance is used, a neighborhood of a point has a spherical shape; when Manhattan distance is used, the shape is rectangular. For simplicity of the presentation, in our examples we will refer to Euclidean distance, although our approach is suitable for any distance metric. Below, we recall definitions of a density based cluster and related notions after [3].

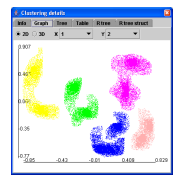
Eps-neighborhood of a point p (denoted by  $N_{Eps}(p)$ ) is defined as the set of points q in dataset D that are distant from p by no more than Eps; that is,  $N_{Eps}(p) = \{q \in D \mid distance(p,q) \leq Eps\}$ .

A point p is defined as a *core point* if its *Eps-neighborhood* contains at least MinPts points; that is, if  $|N_{Eps}(p)| \ge MinPts$ .

A point p is defined as *directly density-reachable* from a point q with respect to Eps and MinPts if  $p \in N_{Eps}(q)$  and q is a core point.



**Fig. 1.** Density-reachability of points (MinPts = 6)



**Fig. 2.** Sample result of clustering with DBSCAN

A point p is defined as *density-reachable* from a point q with respect to Eps and MinPts if there is a sequence of points  $p_1, ..., p_n$  such that  $p_1 = q$ ,  $p_n = p$  and  $p_{i+1}$  is directly density-reachable from  $p_i$ , i = 1..n-1.

**Example 1.** Let MinPts = 6. Point r in Figure 1 has 6 neighbors (including itself) in its neighborhood  $N_{Eps}(r)$ , so it is a core point. Point p has 2 neighbors in  $N_{Eps}(p)$ , so it is not a core point. Point p, however, belongs to  $N_{Eps}(r)$ , so it is directly density-reachable from r. To the contrary r is not directly density-reachable from p despite r belongs to  $N_{Eps}(p)$ . Point p in Figure 1 is density-reachable from point o, since there is a point (e.g. point r) such that p is directly density-reachable from it and it is directly density-reachable from o. Please note that p, which is density-reachable from core point o, is not a core point.  $\square$ 

A point p is defined as a *border point* if it is not a core point, but is density-reachable from some core point. Hence, a point is a border one if it is not a core point, but belongs to the *Eps-neighborhood* of some core point.

Let C(o) determine all points in D that are density-reachable from point o. Clearly, if o is not a core point, then C(o) is empty. In Figure 1, points p and q are density-reachable from core point o. Hence, p and q belong to C(o).

A *cluster*<sup>1</sup> is defined as a non-empty set of all points in D that are density-reachable from a core point. Hence, each C(p) is a cluster provided p is a core point. Interestingly, if p and q are core points belonging to the same cluster, then C(p) = C(q); that is, both points determine the same cluster [3]. Thus, a core point p belongs to exactly one cluster, namely to C(p). We note, however, that a border point may belong to more than one cluster.

*Noise* is defined as the set of all points in D that do not belong to any cluster; that is, the set of all points in D that are not density-reachable from any core point. Hence, each point that is neither a core point, nor border one, constitutes noise.

Fig. 2 presents the results of clustering with DBSCAN for a sample dataset.

# **3** Using the Triangle Inequality for Efficient Determination of Eps-Neighborhoods

Let us start with recalling the triangle inequality property:

Property 1. (Triangle inequality property). For any three points p, q, r:

 $distance(p,r) \le distance(p,q) + distance(q,r)$ 

Property 2 presents its equivalent form, which is more suitable for further considerations.

**Property 2.** (Triangle inequality property). For any three points p, q, r:

 $distance(p,q) \ge distance(p,r) - distance(q,r)$ .

 $<sup>^1</sup>$  This definition differs from the original one provided in [3]. However, it is equivalent to the original one by Lemma 1 in [3], and is more suitable for our presentation.

**Lemma 1.** Let D be a set of points. For any two points p, q in D and any point r:

```
distance(p,r) - distance(q,r) > Eps \implies q \notin N_{Eps}(p) \land p \notin N_{Eps}(q).
```

**Proof.** Let distance(p,r) - distance(q,r) > Eps (\*). By Property 2,  $distance(p,q) \ge distance(p,r) - distance(q,r)$  (\*\*). By (\*) and (\*\*), distance(p,q) > Eps, and distance(q,p) = distance(p,q). Hence,  $q \notin N_{Eps}(p)$  and  $p \notin N_{Eps}(q)$ .  $\square$ 

By Lemma 1, if we know that the difference of distances of two points p and q to some point r is greater than Eps, we are able to conclude that  $q \notin N_{Eps}(p)$  without calculating the actual distance between p and q. Theorem 1 is our proposal of effective determination of points that do not belong to Eps-neighborhood of a given point p.

**Theorem 1.** Let r be any point and D be a set of points ordered in a non-decreasing way with respect to their distances to r. Let p be any point in D,  $q_f$  be a point following point p in D such that  $distance(q_f,r) - distance(p,r) > Eps$ , and  $q_b$  be a point preceding point p in D such that  $distance(p,r) - distance(q_b,r) > Eps$ . Then:

- a)  $q_f$  and all points following  $q_f$  in D do not belong to  $N_{Eps}(p)$ .
- b)  $q_b$  and all points preceding  $q_b$  in D do not belong to  $N_{Eps}(p)$ .

**Proof.** Let r be any point and D be a set of points ordered in a non-decreasing way with respect to their distances to r.

- a) Let p be any point in D,  $q_f$  be a point following point p in D such that  $distance(q_f,r)-distance(p,r)>Eps$  (\*), and s be either point  $q_f$  or any point following  $q_f$  in D. Then  $distance(s,r)\geq distance(q_f,r)$  (\*\*). By (\*) and (\*\*), distance(s,r)-distance(p,r)>Eps. Thus, by Lemma 1,  $s\not\in N_{Eps}(p)$ .
- b) The proof is analogous to the proof of Theorem 1a).  $\Box$

**Corollary 1.** Let r be any point and D be a set of points ordered in a non-decreasing way with respect to their distances to r. Let p be any point in D,  $q_f$  be the first point following point p in D such that  $distance(q_f,r) - distance(p,r) > Eps$ , and  $q_b$  be the first point preceding point p in D such that  $distance(p,r) - distance(q_b,r) > Eps$ . Then, only the points that follow  $q_b$  in D and precede  $q_f$  in D have a chance to belong to  $N_{Eps}(p)$ , and p certainly belongs to  $N_{Eps}(p)$ .

**Example 2.** Let r be a point (0,0). Figure 3 shows sample set D of two dimensional points. Table 1 illustrates the same set D ordered in a non-decreasing way with respect to the distance of its points to point r. Let us consider the determination of the Eps-neighborhood of point p = F, where Eps = 0.5, by means of Corollary 1. We note that distance(F,r) = 3.2, the first point  $q_f$  following point F in D such that distance( $q_f$ ,r) – distance(F,r) > Eps is point C (distance(C,r) – distance(F,r) = 4.5 – 3.2 = 1.3 > Eps), and the first point  $q_b$  preceding point p in D such that distance(F,r) – distance( $q_b$ ,r) > Eps is G (distance(F,r) – distance(G,r) = 3.2 – 2.4 = 0.8 > Eps). By Corollary 1, only the points that follow G and precede C in D (here, points F and H) may belong to  $N_{Eps}(F)$ . Clearly,  $F \in N_{Eps}(F)$ . Hence, H is the only point for which it is necessary to calculate its actual distance to F in order to determine  $N_{Eps}(F)$  properly. □

In the sequel, a point r to which the distances of all points in D have been determined will be called a *reference point*.

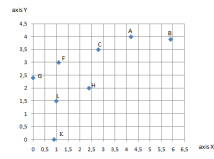


Fig. 3. Set of points D

**Table. 1.** Ordered set of points D from Fig. 3 with their distance to reference point r(0,0)

Q	Χ	Υ	distance(q,r)
K	0,9	0,0	0,9
L	1,0	1,5	1,8
G	0,0	2,4	2,4
Н	2,4	2,0	3,1
F	1,1	3,0	3,2
С	2,8	3,5	4,5
Α	4,2	4,0	5,8
В	5,9	3,9	7,1

# 4 New Algorithms: TI-DBSCAN and TI-DBSCAN-REF

In this section, we propose a new clustering algorithm called TI-DBSCAN and its version TI-DBSCAN-REF. The result of clustering of core points and identifying of noise by our algorithms is the same as the one produced by the DBSCAN algorithm [3]. The border points may be assigned to different clusters<sup>2</sup>. In our algorithms we use Corollary 1 for efficient determination of the Eps-neighborhoods of points. In addition, we adopt the solution from [6] that consists in removing a point from the analyzed set D as soon as it is found to be a core point. Here, we remove each analyzed point, even if it not a core point. Let us start with the description of TI-DBSCAN.

### Notation for TI-DBSCAN

- D the set of points that is subject to clustering;
- Eps the radius of the point neighborhood;
- MinPts the required minimal number of points MinPts within Eps-neighborhood;
- r a reference point assumed to be fixed, e.g. to the point with all coordinates equal to 0 or minimal values in the domains of all coordinates;
- fields of any point p in D:
  - p.ClusterId label of a cluster to which p belongs; initially assigned the UNCLASSIFIED label;
  - o p.dist the distance of point p to reference point r;
  - p.NeighborsNo the number of neighbors of p already found; initially assigned 1 to indicate that a point itself belongs to its own Eps-neighborhood;
  - Border the information about neighbors of p that turned out non-core points for which it is not clear temporary if they are noise ones or border ones; initially assigned an empty set;
- D' the result of clustering of D (initially an empty set);

 $<sup>^2</sup>$  Although a border point may belong to many clusters, DBSCAN assigns it arbitrarily only to one of them, so does TI-DBSCAN. It is easy to modify the algorithms so that border points are assigned to all including clusters.

TI-DBSCAN takes as an input a set of points D, a radius Eps and a threshold MinPts. Its output, i.e. clustered points, shall be stored in D'. A reference point r is assumed to be fixed, e.g. to the point with all coordinates equal to 0 or minimal values in the domains of all coordinates. With each point p in D, there are associated the following fields: ClusterId, dist, NeighborsNo, and Border.

TI-DBSCAN starts with the initialization of D' and the fields of all points in D. Then it sorts all points in D in a non-decreasing way w.r.t. their distance to reference point r. Next it generates a label for the first cluster to be found. Then it scans point after point in D. For each scanned point p, the TI-ExpandCluster function (described later) is called. If p is a core point, then the function assigns the current cluster's label to all points in C(p), moves them from D to D', and TI-DBSCAN generates a next label for a new cluster to be created. Otherwise, point p is assigned label NOISE and is moved from D to D'. After all points in D are scanned, each point is assigned either to a respective cluster identified by ClusterId or is identified as noise.

```
Algorithm TI-DBSCAN(set of points D, Eps, MinPts);
/* assert: r denotes a reference point
D' = empty set of points;
for each point p in set D do
    p.ClusterId = UNCLASSIFIED;
    p.dist = Distance(p,r); p.NeighborsNo = 1; p.Border = \emptyset
endfor
sort all points in D non-decreasingly w.r.t. field dist;
ClusterId = label of first cluster;
{f for} each point p in the ordered set D starting from
    the first point until last point in D do
     if TI-ExpandCluster(D, D', p, ClusterId, Eps, MinPts) then
         ClusterId = NextId(ClusterId)
     endif
endfor
                                     // D' is a clustered set of points
return D'
```

The TI-ExpandCluster function starts with calling TI-Neighborhood function (decribed later) to determine Eps-neighborhood of a given point p in, possibly reduced, set D (more precisely, TI-Neighborhood determines  $N_{Eps}(p) \setminus \{p\}$  in D) and stores it in the seeds variable. Clearly,  $N_{Eps}(p)$  determined in a reduced set D will not contain the neighboring points that were already moved from D to D'. In order to determine the real size of  $N_{Eps}(p)$  in the original, non-reduced D, the auxiliary NeighborsNo field of point p is used. Whenever, a point p is moved from D to D', the NeighborsNo field of each of its neighboring points in D is incremented. As a result, the sum of the size of  $N_{Eps}(p)$  found in the reduced D and p.NeighborsNo equals the size of  $N_{Eps}(p)$  in the original, non-reduced set D.

If p is found not to be a core point, it is temporary labeled as a noise point, NeighborsNo field of each of its neighboring points in D is incremented, the information about p is added to the Border field of each of its neighboring points in D, p itself is moved from D to D', and the function reports failure of expanding a cluster.

Otherwise, the examined point is a core point and all points that are density-reachable from it will constitute a cluster. First, all points in the Eps-neighborhood of the analyzed point are assigned a label (ClId) of the currently built cluster and their NeighborsNo fields are incremented. Next all non-core points indicated by the

p.Border, which were stored in D', are found to be border points, and are assigned cluster label ClId, p.Border is cleared, and p is moved from D to D'. Now, each core point in seeds further extends the seeds collection with the points in its Epsneighborhood that are still unclassified. After processing a seed point, it is deleted from seeds. The function ends when all points found as cluster seeds are processed.

Note that TI-ExpandCluster calculates Eps-neighborhood for each point only once.

```
\textbf{function} \  \, \texttt{TI-ExpandCluster}(\textbf{var} \  \, \texttt{D, var} \  \, \texttt{D', point p, ClId, Eps, MinPts})
/* assert: D is ordered in a non-decreasing way w.r.t.
  distances of points in D from the reference point r.
/* assert: TI-Neighborhood does not return p.
seeds = TI-Neighborhood(D, p, Eps);
p.NeighborsNo = p.NeighborsNo + |seeds|;
                                                         // including p itself
if p.NeighborsNo < MinPts then</pre>
                                      // p is either noise or a border point
    p.ClusterId = NOISE;
    for each point q in seeds do
         append p to q.Border; q.NeighborsNo = q.NeighborsNo + 1
    p.Border = \emptyset; move p from D to D';
                                               // D' stores analyzed points
    return FALSE
    p.ClusterId = ClId;
    \textbf{for} \ \texttt{each} \ \texttt{point} \ \texttt{q} \ \texttt{in} \ \texttt{seeds} \ \textbf{do}
         q.ClusterId = ClId; q.NeighborsNo = q.NeighborsNo + 1
    endfor
    \textbf{for} \ \texttt{each point q in p.Border} \ \textbf{do}
        D'.q.ClusterId = ClId;
                                     //assign cluster id to q in D'
    endfor
    p.Border = \emptyset; move p from D to D';
                                                 // D' stores analyzed points
    while |seeds| > 0 do
        curPoint = first point in seeds;
         curSeeds = TI-Neighborhood(D, curPoint, Eps);
         curPoint.NeighborsNo = curPoint.NeighborsNo + |curSeeds|;
         if curPoint.NeighborsNo < MinPts then //curPoint is border point</pre>
             for each point q in curSeeds do
                  q.NeighborsNo = q.NeighborsNo + 1
             endfor
         else
                                                   // curPoint is a core point
             for each point q in curSeeds do
                  q.NeighborsNo = q.NeighborsNo + 1
                  if q.ClusterId = UNCLASSIFIED then
                       q.ClusterId = ClId;
                      move q from curSeeds to seeds
                  else
                      delete q from curSeeds
                  endif
             endfor
             for each point q in curPoint.Border do
                   D'.q.ClusterId = ClId; //assign cluster id to q in D'
             endfor
         endif
         curPoint.Border = \emptyset; move curPoint from D to D';
         delete curPoint from seeds
    endwhile
    return TRUE
endif
```

The TI-Neighborhood function takes the ordered point set D, point p in D, and Eps as input parameters. It returns  $N_{Eps}(p)\setminus\{p\}$  as the set theoretical union of the point sets found by the TI-Backward-Neighborhood function and the TI-Forward-Neighborhood function. TI-Backward-Neighborhood examines points preceding currently analyzed point p, for which Eps-neighborhood is to be determined. The function applies Lemma 1 to identify first point, say q<sub>b</sub>, preceding p in D such that distance(p,r) distance $(q_b,r)$  > Eps. All points preceding point  $q_b$  in D are not checked at all, since they are guaranteed not to belong to  $N_{Eps}(p)$  (by Theorem 1). The points that precede p and, at the same time, follow  $q_b$  in D have a chance to belong to  $N_{Eps}(p)$ . For these points, it is necessary to calculate their actual distance to p (When using the Euclidean distance metric, the functions may apply the square of Distance and the square of Eps for efficiency reasons). The TI-Backward-Neighborhood function returns all points preceding p in D with the distance to p not exceeding Eps. The TI-Forward-Neighborhood function is analogous to TI-Backward-Neighborhood. Unlike TI-Backward-Neighborhood, TI-Forward-Neighborhood examines points following currently analyzed point p, for which Eps-neighborhood is to be determined. The TI-Forward-Neighborhood function returns all points following p in D with the distance to p not exceeding Eps.

```
function TI-Neighborhood(D, point p, Eps)
return TI-Backward-Neighborhood(D, p, Eps) ∪
        TI-Forward-Neighborhood(D, p, Eps)
function TI-Backward-Neighborhood(D, point p, Eps)
/* assert: D is ordered non-decreasingly w.r.t. dist */
seeds = \{\}:
backwardThreshold = p.dist - Eps;
for each point q in the ordered set D starting from
    the point immediately preceding point p until
    the first point in D do
    if q.dist < backwardThreshold then</pre>
                                                    // p.dist - q.dist > Eps?
         break;
    endif
    \textbf{if} \ \texttt{Distance} \ (\texttt{q, p}) \ \leq \ \texttt{Eps} \ \textbf{then} \ \texttt{append} \ \texttt{q} \ \texttt{to} \ \texttt{seeds} \ \textbf{endif}
endfor
return seeds
function TI-Forward-Neighborhood(D, point p, Eps)
/* assert: D is ordered non-decreasingly w.r.t. dist */
seeds = \{\};
forwardThreshold = Eps + p.dist;
for each point q in the ordered set D starting from
    the point immediately following point p until
```

// q.dist - p.dist > Eps?

the last point in D do

endif

return seeds

if q.dist > forwardThreshold then

if Distance(q, p)  $\leq$  Eps then append q to seeds endif

Except for TI-DBSCAN, we have also implemented its variant TI-DBSCAN-REF [5] that uses many reference points instead of one for estimating the distance among pairs of points. Additional reference points are used only when the basic reference point according to which the points in D are sorted is not sufficient to estimate if a given point q belongs to Eps-neighborhood of another point p. The estimation of the distance between q and p by means of an additional reference point is based on Lemma 1. The actual distance between the two points q and p is calculated only when all reference points are not sufficient to estimate if  $q \in N_{Eps}(p)$ .

In fact, TI-DBSCAN-REF was obtained from TI-DBSCAN by introducing the following in the functions: TI-DBSCAN, TI-Backward-Neighborhood and TI-Forward-Neighborhood. Below we provide the modified versions of these functions. The introduced changes are highlighted in the code.

In our many-reference-points version of TI-DBSCAN, all reference points are stored in array RefPoints. In addition, each point p in set D of points to be clustered will be associated with the array Dists, storing distances between p and referenced points stored in RefPoints. The first point in Dists (that is, Dists[1]) plays the same role as the field dist in the one-reference-point version of TI-DBSCAN. Consequently, all points in D will be sorted in a non-decreasing way with respect to their distances Dists[1] to the first reference point RefPoints[1].

```
Algorithm TI-DBSCAN-REF(set of points D, Eps, MinPts);
store reference points in array RefPoints;
ClusterId = label of first cluster;
for each point p in set D do
    p.ClusterId = UNCLASSIFIED
    for i = 1 to |RefPoints| do
        p.Dists[i] = Distance(p, RefPoints[i]);
    endfor
    p.NeighborsNo = 1; p.Border = \emptyset
endfor
sort all points in D non-decreasingly w.r.t. attribute Dists[1];
for each point p in the ordered set D starting from
    the first point until last point in D do
    \textbf{if} \ \texttt{TI-ExpandCluster}(\texttt{D, D', ClusterId, Eps, MinPts}) \ \textbf{then}
         ClusterId = NextId(ClusterId)
endfor
                                    // D' is a clustered set of points
return D'
```

The many-reference-points version of TI-Backward-Neighborhood function differs from one-reference-point version only in treating the points in D that were not excluded from the analysis carried out by means of triangle inequality property applied to the first reference point RefPoints[1], a given point p, and a point q being a candidate for a neighbor of point p; that is based on Dists[1] fields of p and q. In one-reference-point version of TI-Backward-Neighborhood, the neighborhood between point p and a non-excluded point q requires the calculation of the actual distance between the two points. In many-reference-points version, this calculation is deferred; namely, the actual distance calculation between p and q is carried out only when the usage of triangle inequality property with respect to all remaining reference points in RefPoints does not invalidate q as a neighbor of p (based on respective Dists[i] for p

and  $q, i \ge 2$ ). Otherwise, q is find not to be a neighbor of p without knowing the actual distance between the two points.

```
function TI-Backward-Neighborhood(D, point p, Eps, MinPts)
/* assert: D is ordered non-decreasingly w.r.t. Dists[1] */
seeds = \{\};
backwardThreshold = p.Dists[1] - Eps
for each point q in the ordered set D starting from
    the point immediately preceding point p until
    the first point in D do
     /* p.Dists[1] - q.Dists[1] > Eps? */
    if q.Dists[1] < backwardThreshold then</pre>
       break;
    endif
    candidateNeighbor = true;
    i = 2;
    while candidateNeighbor and (i ≤ |p.Dists|) do
   if |q.Dists[i] - p.Dists[i]| > Eps then
             candidateNeighbor = false
        else
        endif
    endwhile
    if candidateNeighbor then
        if Distance(q, p) \le Eps then append q to seeds endif
endfor
return seeds
```

The many-reference-points version of the TI-Forward-Neighborhood function applies analogous adaptation as in the case of TI-Backward-Neighborhood.

```
\textbf{function} \; \texttt{TI-Forward-Neighborhood(D, point p, Eps, MinPts)}
/* assert: D is ordered non-decreasingly w.r.t. Dists[1]
seeds = {};
forwardThreshold = Eps + p.Dists[1]
for each point q in the ordered set D starting from
    the point immediately following point p until
    the last point in D \boldsymbol{do}
    /* q.Dists[1] - p.Dists[1] > Eps? */
    if q.Dists[1] > forwardThreshold then
         break;
    endif
    candidateNeighbor = true;
    i = 2;
    while candidateNeighbor and (i ≤ |p.Dists|) do
   if |q.Dists[i] - p.Dists[i]| > Eps then
             candidateNeighbor = false
         else
             i = i + 1
         endif
    endwhile
    if candidateNeighbor then
         if Distance(q, p) \le Eps then append q to seeds endif
    endif
endfor
```

# **5 Performance Evaluation**

We have examined the following versions of the DBSCAN algorithm: DBSCAN-RTree which uses R-Tree as an index, TI-DBSCAN using one reference point, TI-DBSCAN-Ref using as many points as the number of dimensions of a clustered dataset and TI-DBSCAN-Ref-Mink taking advantage of Minkowski distance as well as many reference points.

In this section, we report the results of our experimental evaluation of the above mentioned algorithms. In the experiments, we used a number of datasets (and/or their subsamples) of different cardinality and dimensionality. In particular, we used widely known datasets such as: birch [9], SEQUOIA 2000 [8], and kddcup 98 [10] as well as datasets generated automatically (random) or manually.

**Table 2.** Datasets used in experiments. Notation: dim. – number of point's dimensions, card. – number of points in a dataset.

No	Name	dim.	card.		
1	sequoia_2000_d2_1252	2	1252		
2	sequoia_2000_d2_2503	2	2503		
3	sequoia_2000_d2_3910	2	3910		
4	sequoia_2000_d2_5213	2	5213		
5	sequoia_2000_d2_6256	2	6256		
6	sequoia_2000_d2_62556	2			
7	manual_d2_2658	2	2658		
8	manual_d2_14453	2	14453		
9	random_d3_50000	3	50000		
10	random_d10_10000	10	10000		
11	random_d10_20000	10	20000		
12	random_d10_50000	10	50000		
13	random_d20_500	20	500		
14	random_d40_500	40	500		
15	random_d200_500	200	500		
16	random_d200_1000	200	1000		
17	birch_d2_100000	2	100000		
18	random_d100_1000	100	1000		
19	random_d50_10000	50	10000		
20	random_d100_10000	100	10000		
21	random_d50_20000	50	20000		
22	random_d100_20000	100	20000		
23	random_d5_100000	5	100000		
24	kddcup_98_d56_56000	56	56000		

The run times of clustering with TI-DBSCAN, TI-DBSCAN-Ref, TI-DBSCAN-REF-Mink, and DBSCAN using R-Tree as an index are presented in Table 3.

**Table 3.** Datasets used in experiments and run times (in milliseconds) of examined algorithms. Notation: dim. – number of point's dimensions, card. – number of points in a dataset, prep. – time of sorting of points, ind. – time of building of an index, clust. – clustering, "-" – results not available within at least 12 hours or due to memory limitations.

			TI-DBSCAN-REF-Mink		TI-DBSCAN-Ref		TI-DBSCAN		DBSCAN-RTree	
No dim.		card.	minPts = 4, Eps = 10, m = 1		minPts = 4, Eps = 10		minPts = 4, Eps = 10		minPts = 4, Eps = 10	
			prep.	clust.	prep.	clust.	prep.	clust.	ind.	clust.
1	2	1252	11	12	12	10	11	5	1760	959
2	2	2503	24	13	22	8	22	8	482	1469
3	2	3910	41	21	41	18	43	17	632	2929
4	2	5213	83	24	80	15	82	12	928	3466
5	2	6256	97	26	100	33	94	19	1087	4201
6	2	62556	21897	180	21326	160	21775	95	8497	53384
7	2	2658	27	36	26	92	27	98	606	2536
8	2	14453	823	738	645	1352	802	1535	2629	15381
9	3	50000	10017	941	10284	7964	10706	16772	10463	76951
10	10	10000	245	116	246	267	231	2072	6559	124496
11	10	20000	886	265	889	1000	931	8347	12962	378147
12	10	50000	7902	697	13932	6265	7459	53140	31403	1291729
13	20	500	11	24	10	14	10	13	534	1604
14	40	500	10	40	9	35	10	23	1297	3033
15	200	500	14	821	14	710	14	114	-	-
16	200	1000	21	1701	25	1450	25	424	-	-
17	2	100000	73877	251	78341	153	103725	97	21150	66974
18	100	1000	20	399	21	412	23	219	10560	24399
19	50	10000	274	1202	285	1198	305	10043	44553	1388936
20	100	10000	300	4646	297	4051	342	19248	133568	2511244
21	50	20000	973	2607	952	3134	1163	41536	93973	6189963
22	100	20000	1062	8496	1065	8658	1048	81828	258368	10725222
23	5	100000	66132	1237	77092	22356	70557	118964	30779	353544
24	56	56000	69830	16712	130575	53736	76028	240818	-	-

As follows from Table 3, TI-DBSCAN and TI-DBSCAN-Ref are more efficient than DBSCAN with R-Tree even up to 600 times. TI-DBSCAN-Ref tends to be faster than TI-DBSCAN for large high dimensional datasets. For small low dimensional datasets, TI-DBSCAN tends to be faster than TI-DBSCAN-Ref. Additionally, the version using the Minkowski distance, is even more efficient than TI-DBSCAN-Ref for larger datasets.

# **6 Conclusions**

In the paper, we have proposed two versions of our new algorithm: TI-DBSCAN and TI-DBSCAN-Ref that produce the same results as DBSCAN, but use the triangle inequality to speed up the clustering process. TI-DBSCAN uses only one reference point, while TI-DBSCAN-Ref uses many reference points. As follows from the experiments, both versions of our algorithm are much more efficient than the original DBSCAN algorithm, which is supported by a spatial index. Unlike DBSCAN, TI-DBSCAN and TI-DBSCAN-Ref enable efficient clustering of high-dimensional data. Moreover, for larger datasets, the version of TI-DBSCAN using the Minkowski distance (TI-DBSCAN-Ref-Mink) is more efficient then TI-DBSCAN-Ref. The usage of many reference points and the Minkowski distance is particularly useful in the case of large high dimensional datasets.

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