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Improved density peak clustering for large datasets

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Abstract

Clustering is the usual way of classifying data when there is no a priori knowledge, especially about the number of classes. Within the frame of big data analysis, the computational effort needed to perform the clustering task may become prohibitive and motivated the construction of several algorithms or the adaptation of existing
ones, as the well known K-means algorithm [12]. Recently, Rodriguez and Laio [17] proposed an algorithm that clusters efficiently by fast searching local density peaks that are sufficiently distant one from the others. However it is able to work on small datasets only and is highly sensitive to the value of tunable parameters. In this paper we propose Improved Density Peak Clustering (IDPC), a new algorithm designed for large datasets based on [17] which corrects the shortcomings mentioned above. Thanks to our Cover Map (CM) procedure iterated with a decreasing locally-adaptive window (ICMDW), we are able to build both a localisation map and a multidimensional density map. The nature of the density map, which fits perfectly with the approach of [17], allows us to compute the different steps with much less operations. It carries unsensitive parameters, supports last improvements on cluster centers selection and potentially allows new improvements.

Keywords: Clustering algorithm; density-based clustering; large datasets;

1 Introduction

Clustering is a commonly used tool that aims to identify similar samples in a dataset. It can be viewed both from the machine learning and the statistics point of view, with classical algorithms pertaining to each domain. When considered from the algorithmic standpoint, the complexity of the clustering problem is known to be NP hard, even for the usual K-means, when the number of clusters is not fixed.

Presented in [17], the innovative fast density peak detection (FDPC) algorithm is able to cope with non-convex clusters and gives a convenient decision rule to find the correct number of clusters. As in the mean shift algorithm [3], clusters centers are defined as local density maxima. More specifically, a cluster center is defined as a point surrounded by neighbours with lower local density and away from any other local maxima. In order to detect them, a distance $d_c$ is first selected and is used to define neighborhood. Densities $\rho_i$ are calculated at each data point $x_i$ by counting the number of sample points closer than
$d_c$ to $x_i$ (1). Next, for each data point, the nearest point with a higher density is found and the distance $\delta_i$ separating them is recorded (2). Cluster centers are then defined as the points $x_i$ that have both high density $\rho_i$ and high distance $\delta_i$. Other points receive the same label than their nearest neighbor of higher density (3).

This innovative method has promising results but shows some limitations. In step (1), the distances between all sample points are calculated, leading to an $O(N^2)$ complexity and memory footprint, not to mention other steps. This issue limits it to small datasets. Furthermore, choosing a good threshold distance $d_c$ is non-trivial and critical since it has a major impact on the final clustering.

Our new algorithm aims to overcome the drawbacks mentioned above. The general idea of IDPC is to start by the construction of a density map through the procedure Iterated Cover Map with a Decreasing Window (ICMDW), reducing the dimension $N$ of the problem, and then to use the localisation information it provides to construct a very fast version of steps (2) and (3).

In the second Section we review the work related to [17]. In the third Section we detail the implementation of the ICMDW algorithm. In the fourth Section we explain how the results provided by ICMDW are used to greatly reduce the computations in cluster centroid selection and labelisation. And finally, we present experimental results in the fifth Section.

2 Related work

2.1 Fast Density Peak Clustering

Let $(x_i)_{i=1,...,N}$ be the sample points. In the first step of FDPC the Euclidean distances $d_{ij}$ between each data points are computed. Then a distance $d_c$ is chosen to finally compute, for each $x_i$, the number of data points that are closer than $d_c$ to the point $x_i$. Local densities
are then obtained according to the formula:

\[ \rho_i := \rho(x_i) = \sum_{j=1}^{N} (1_{R}(d_{ij} - d_c)). \]

Please note that the expression is of the form \( \sum_{j=1}^{N} K(x_i - x_j) \), where \( K \) is the so-called kernel function, that reduces here to the characteristic function of a ball \( 1_{B(0,d_c)} \).

In the second step, the minimum distance between the point \( x_i \) and any other point with higher density is calculated as \( \delta_i = \min_{j: \rho_j > \rho_i} d_{ij} \). Finally, cluster centers are found as outliers in the decision graph formed by \( (\delta, \rho) \), where outliers are the points sharing anomalously large density-delta values. Conventionally, for the point \( x_k \) with highest density, \( \delta_k = \max_j (d_{ij}) \).

Finally, other points are assigned to the same cluster than their Nearest Point with Higher Density (NHPD). Its algorithmic is detailed in Algorithm 1.

Many researchers worked on this algorithm in order to improve and/or use it on a specific problem. It has been used successfully on clustering of electricity consumption behaviour [22], text clustering [10], unsupervised acoustic subword units discovery pompage [25], batch process modelling and on-line monitoring [16], medical data [7] and it has been

---

**Algorithm 1 FDPC**

```
procedure FDPC(d_c)
    for all \((i,j) \in \{1,...,N\}^2\) do
        \(d_{ij} \leftarrow \text{distance}(x_i, x_j)\)
    end for
    for all \(i \in \{1,...,N\}\) do
        \(\rho_i \leftarrow \sum_{j=1}^{N} (1_{x<0} (d_{ij} - d_c))\)
    end for
    for all \(j \in \{1,...,N\}\) do
        \(\text{nhpd}_j \leftarrow \arg\min_{k: \rho(x_k) > \rho(x_j)} (d_{jk})\)
        \(\delta_j \leftarrow d_{j,\text{nhpd}_j}\)
    end for
    Cluster centers \(\leftarrow\) outliers in decision graph
    \(\forall j,\text{ Label}_j \leftarrow \text{Label}_{\text{nhpd}_j}\)
end procedure
```
slightly modified to work on image segmentation [1, 2, 18], outliers detection [4] and hyper
spectral band selection [6]. K-means and FDPC have been compared in [15] and FDPC
has shown more accurate than K-means.

However, FDPC has proved to be slower than K-means and suffers from three other
shortcomings:

- First of all, due to its inherent computational complexity, it cannot handle large data
  sets.

- Secondly, the choice of $d_c$ has to be made empirically whereas it highly impacts every
  steps of the clustering algorithm.

- Finally, no decision rule is given to determine the outliers in the decision graph,
  leading to a manual or poor automatic selection of the cluster centers.

2.2 Improvements of FCDP

To overcome the limitations underlined above, a lot of work was done to derive improved
versions of the original algorithm. Most of them deals only with one issue, due to the
application:

**Ability to handle large datasets**

An extension of FDPC [9] has been proposed in order to manage large taxi fleet datasets.
Data sample is first projected to a density image which is processed to obtain the densities
$\rho_i$ and the contours of clusters. This method works well on large datasets, but for 2-
dimensional data only.

**Choice of $d_c$**

The baseline choice of $d_c$ is to take the one that produce an average density count equal
to 2% of $N (\bar{\rho} = 2\%N)$[4]. A non-trivial entropy-based choice of $d_c$ has been brought in
[20], but it is computationally difficult to perform, even more in large scale contexts. There
exist two methods that replace the density estimation (1) by a non-parametric one, avoiding the choice of \( d_c \). In [21] the local density is estimated with a non-parametric multivariate kernel estimator. For each point \( x_i \), the local density is calculated throughout the formula

\[
\rho_i = \frac{1}{n \prod_{l} h_l} \sum_{j=1}^{n} K \left( \frac{x_{i1} - x_{j1}}{h_1}, ..., \frac{x_{id} - x_{jd}}{h_d} \right),
\]

where \( d \) is the dimension, \( K \) is a Gaussian kernel and \( \{h_1, ..., h_d\} \) are the bandwidths which are automatically and locally chosen. Moreover an automatic cluster centroid selection method is also developed through maximizing an average silhouette index. In [14], density estimation is done with an another non-parametric density estimator, based on the heat equation, which is more accurate but computationally expensive.

**Cluster centers selection**

In [11] data points are arranged according to their \( \gamma_i = \delta_i \cdot \rho_i \) values in descending order and the first \( m \) points are selected. Then centers are the points that possess a higher \( \gamma_i \) than the turning-point defined as

\[
\arg\max_{i \in \{1, ..., m\}} \left( \frac{k_i^1}{k_i^{n-1}} \right), \quad \text{where} \quad k_i^n = \frac{\gamma_i + n - \gamma_i}{n}.
\]

Regarding [13], cluster centers are supposed to possess large \( \delta_i \) (\( \delta_i > 2 \text{Var}(\delta) \) in [13], \( \delta_i > 3 \text{Var}(\delta) \) in [4]) and large \( \rho_i \) (\( \rho_i > \text{mean}(\rho) \)). Furthermore, a merge step is added: two clusters are merged if it exists a point \( x \) belonging to one cluster and a point \( y \) belonging to the other one that are closer than \( d_c \). [23] also merges recursively the clusters based on their relative inter-connectivity and relative closeness.

An another \( d_c \)-based algorithm is presented in [8]. The concepts of DBSCAN [5] (density reachable, core objects..) are coupled with the divide-and-conquer strategy to produce a better cluster centers selection. Potential centroids obtained by the decision graph are
tested recursively and ignored if there is density-reachability between it and any other cluster centroid.

A $k$-nearest neighbors (KNN) based algorithm [19] is proposed as a rigorous way to detect cluster centers automatically. KNN is used to define $\rho_i = \frac{1}{\sum_{j=1}^{K} d_{ij}}$. The product $\gamma = \rho \cdot \delta$ is proven to follow a fat tailed distribution and a statistical test is proposed to find its outliers, which will define the cluster centers.

Other improvements

In [24], KNN is used to improve the assignment rule of the remaining points, right after the cluster centers detection step. First, outlier points (with a large $\delta_i^K = \max_{i \in \text{KNN}_i}(d_{ij})$) are deleted. Secondly, remaining points are assigned through a restrictive strategy called "strategy 1". Roughly speaking, the label of a centroid $x_c$ is propagated to the unlabelled points closer than $\sum_{j \in \text{KNN}_c} d_{cj} / K$ and those newly labelled points propagates recursively the label by the same way. Unassigned points by the strategy 1 are managed by the second strategy. The key aspect of strategy 2 is to learn the probability $p_i^c$ that a point $x_i$ belongs to cluster $c$, then assign the point $i$ to its most similar cluster $c$ with the highest value of $p_i^c$.

In [26], a derivative of FDPC that can manage uneven density datasets is proposed. The differences occurs for densities $\rho_i$ and distances $\delta_i$ that are replaced by quantities with a similar behavior. The notion of mutual attraction of two points is introduced and is given by

$$f_{ij} = \min(f_{i \rightarrow j}, f_{j \rightarrow i}), \text{ where } f_{i \rightarrow j} = e^{-\frac{d(i,j)^2}{2\sigma^2}}.$$ 

The parameter $\sigma$, controlling the width of the kernel function, is locally calculated by

$$\sigma_i = \frac{\sum_{j \in J} d(i,j)}{\text{Card}(J)}, \text{ where } J = \{j | d(i,j) \leq d_c\}.$$
The densities \( \rho_i \) are then replaced by the influence scores

\[
\rho_i = \frac{\sum_{j \in A} f_{i,j}}{\text{card}(A)}
\]

and the \( \delta_i \) are calculated with

\[
\delta_i = \min_{j: \rho_j > \rho_i} \frac{d(i,j)}{\min(\sigma_i, \sigma_j)}.
\]

3 Building density map and localisation map

In order to address the issue of large databases, instead of evaluating the densities at every single point \( x_i \in N \times P \), we propose to evaluate the densities \( \rho_i \) at a reduced number of points covering the domain of the data. Data sample is randomized and pseudo normalized (normalized but with a small part of the dataset). The densities \( \rho_i \) are calculated with an adaptive windows \( d^g_c \), smaller when the local density is high. To do so, the procedure Cover Map (CM) is iterated with a decreasing window size \( d^g_c \). At each iteration \( g \) the results obtained from the last CM iteration are used to perform a cheaper CM than the one that would be directly performed with \( d^g_c \).

3.1 Procedure Cover Map

For a given fixed window \( d_c \), the Cover Map procedure covers the domain of the dataset with balls of radius equal to \( d_c \).

The first observation is used to construct the first ball \( B_1(x_1, d_c) \). Then for each observation \( x_i \) it checks if the data point belongs to at least one existing ball. If yes, then those balls increase their density count by one. Else, a new ball \( B(x_i, d_c) \) is created (see algorithm 2).

Thanks to this procedure we obtain a subset of points, possessing much fewer points that the original one, with their associated local density \( \rho_i \) covering the whole domain. But it remains the difficult choice of \( d_c \), and the reduced set of balls can still be too large for the second step.
Algorithm 2 Cover Map

1: procedure $CM(d_c)$
2: $S \leftarrow B_1(x_1, d_c)$  \hspace{1cm} \triangleright \text{Create the first ball}$
3: $\rho_1(B_1(x_1, d_c)) \leftarrow 1$
4: for all $x_i, i > 1$ do
5: \hspace{1cm} $\overrightarrow{d} \leftarrow \text{distance}(x_i, S)$
6: \hspace{1cm} $J \leftarrow \{j | \overrightarrow{d}[j] < d_c\}$
7: \hspace{1cm} if $J \neq \emptyset$ then
8: \hspace{2cm} $\forall j \in J, \rho_j(B_j) \leftarrow \rho_j(B_j) + 1$
9: \hspace{1cm} else
10: \hspace{2cm} $S \leftarrow S \cup B_k(x_i, d_c)$  \hspace{1cm} \triangleright \text{Create new ball}$
11: \hspace{2cm} $\rho_k(B_k(x_i, d_c)) \leftarrow 1$
12: \hspace{1cm} end if
13: end for
14: return $\{S, \{\rho_k\}_k\}$
15: end procedure

3.2 Iterative CM procedure with a decreasing window (ICMDW)

A modified version of the CM procedure described above is iterated with a decreasing window $d^g_c = d^{g-1}_c R$, $R \in ]0,1[$. At each consecutive CM iteration it uses the results of the last one in order to reduce the number of distances to be calculated. The modified CM algorithm mainly differs in the construction and use of a family-tree $F_{\text{tree}}$ which holds the localisation information of the balls, allowing much less computations. This algorithm creates a collection of balls with different size $\{B^g_j\}_{g,j}$, the corresponding local densities and the family-tree $F_{\text{tree}}$. The various terms used in the sequel are summarized below.

**Terminology:**

**Terminal ball:** A terminal ball is a ball that contains less than $N_{\text{min}}$ data points.

**Attached:** A ball $B^g(x, d_c \times R)$ is attached to a previous generation ball $B^{g-1}(y, d_c)$ if $d(x, y) < d_c.(R + 1)$.

**Daughters of $B$: $D(B)$,** the daughters of the ball $B$ are the next generation balls attached to it.

**Mother of $B^g(x, d_c)$:** $M(B^g(x, d_c))$, the mother of $B$ is the closest ball of generation $g - 1$
containing $x$.

**Mother of $x$:** When the current generation is $g$, the mother of the data point $x$ is the closest ball of generation $g - 1$ containing $x$. It is denoted by $M(x)$.

**Family-tree:** The family-tree contains the balls created at each generation and their links.

We enforce that if a data point $x_i$ belongs to a ball $B^g$ of generation $g$ then it cannot belong to a ball of next generation $g + 1$ unattached to $B^g$. Formally, if $x_i \in B^g_j$ and $x_i \in B^{g+1}_k$ then $B^{g+1}_k$ is attached to $B^g_j$.

This implies that when we check a new observation at generation $g$ in the CM procedure, we do not need to calculate the distances between it and all existing balls (their number can be very high), but we only need to compute the distances between it and the few balls of generation $g$ attached to the mother ball (generation $g - 1$) of $x_i$.

The first step of ICMDW (see Algorithm 3) is a standard CM iteration with a very large initial $d^1_c$. Then the modified CM algorithm (see Algorithm 4) is iteratively applied with a smaller window $d^2_c = d^1_c \cdot R$, where $R \in ]0, 1[$ is the coefficient which controls the decreasing speed of the window. At a given iteration $g$, the modified CM procedure works as follow. For each $x_i$ it first constructs $J$, the set of indexes of existing balls to which $x_i$ belongs to. But for that purpose, instead of comparing $x_i$ with the whole set $S$ of existing balls, it only searches among the reduced set $s$. $s$ is constructed by loading the mother $m = M(x_i)$ of $x_i$ and then by loading $s = D(m)$, the daughters balls of $m$. $J$ can then be obtained by calculating the distances between $x_i$ and the reduced set of balls $s$ only. If $J \neq \emptyset$, the balls pointed by $J$ increase their density count by one. Else, a new ball $B(x_i, d^g_c)$ is created and $F_{\text{tree}}$ is updated by indicating to which balls $B(x_i, d^g_c)$ is attached. For this update, which happens only when a ball is created, we need to compute the distance between $x_i$ and all the current generation existing balls to retain those closer than $d^g_c \cdot (1 + R)$. Finally, $J$ is stored to be able to load its mother and to know if $x_i$ belongs to terminal balls in the
next CM iteration. After all observations $x_i$ has been processed, the balls with less than $N_{\text{min}}$ points are set terminals and data points that belongs to only terminal balls are frozen (not used in the next iterations), reducing the number of data points processed at next the rounds. The frozen points from the last round are nevertheless used one last time. Indeed, data points frozen at iteration $g$ are used at iteration $g + 1$ to increase the density count of the $\rho_i^{g+1}$, but not to create new balls. This procedure is iterated and stops when there remain only terminal balls.

Density counts are re-evaluated through a last pass of the dataset. To do so, for each point $x_i$, we find the balls of first generation to which $x_i$ belongs to, increase their density count by one, load the daughters of the closer one and repeat the process until $x_i$ doesn’t belong to any of the daughter balls. From those operations we can also obtain $cc_i$, for each data-point $x_i$, the closest and smaller terminal ball containing it (used for the labelisation phase). This re-evaluation is necessary since some balls that are created lately comes after the drawing of many points that should belong to them, implying an artificial small density count. Finally densities are rescaled with respect to the volume of the balls.

In [4] $d_c$ is chosen such as \( \bar{\rho} = 2\%N \). However, here it is the width of the window $d^g_c$ which is indirectly defined by $N_{\text{min}}$. Since we work in a large scale context and since the window is adaptive, we can choose $N_{\text{min}}$ much lower than $2\%N$. From our experience, we propose $N_{\text{min}} = 0.1\%N$ but it can be modified. It can be raised for faster but less accurate results, or parsimoniously lowered for expensive but more accurate results. Note that the accuracy is not of much importance for the cluster centers selection, but is determinant for the labelisation phase.

$R \in [0, 1]$ controls the decreasing speed of the radius. When $R$ is small, the balls will decrease faster implying better performances, but will increase the risks of creating balls that possess low $\rho_i$ while the area may be dense. To overcome this problem $N_{\text{min}}$ can be raised but it will imply a loss in details since many balls will not be deployed. Setting
Algorithm 3 Iterated CM With Decreasing Window

1: **procedure** ICMDW($d_c, N_{\text{min}}, R$)

2: $d^1_c \leftarrow R$

3: $\{B^1_k\}_k, \{\rho^1_k\}_k \leftarrow \text{CM}(d^1_c)$

4: $g \leftarrow 1$

5: **while** it remains at least a non-terminal ball **do**

6: $g \leftarrow g + 1$

7: $d^g_c \leftarrow d^{g-1}_c \cdot R$

8: $\{F_{\text{tree}}, \rho^g_k, S^g, TB^g\} \leftarrow \text{CM}_{\text{modif}}(d_c, N_{\text{min}}, R, F_{\text{tree}}, g, S^{g-1})$

9: **end while**

10: $\triangleright$ Density re-evaluation:

11: **for all** $x_i \in \{1, ..., N\}$ **do**

12: $s \leftarrow \{B^1_k\}_k$

13: $g \leftarrow 1$

14: **while** $J \neq \emptyset$ **do**

15: $\overrightarrow{d} \leftarrow \text{distance}(x_i, s)$

16: $J \leftarrow \{j | \overrightarrow{d}[j] < d^g_c\}$

17: $\forall j \in J, \rho^g_j \leftarrow \rho^g_j + 1$

18: $cc_i \leftarrow B^g_{\overrightarrow{J}[1]}$

19: $g \leftarrow g + 1$

20: $s \leftarrow D(J[1])$

21: **end while**

22: **end for**

23: $\{\rho^g_k\} \leftarrow \{\rho^g_k / \text{Volume}^g\}$

24: **return** $\{F_{\text{tree}}, \rho^g_k, S^g, TB^g, \{cc_i\}_{i \in \{1, ..., N\}}\}$

25: **end procedure**
Algorithm 4 CM modified

1: procedure $CM_{MODIF}(d_c, N_{min}, R, F_{tree}, g)$
2: \hspace{0.5em} $S \leftarrow \emptyset$
3: \hspace{0.5em} for all not frozen $x_i$ do
4: \hspace{1em} $m \leftarrow M(x_i)$ \hspace{1em} \text{\texttt{\texttt{\texttt{\texttt{\triangle}}}} Load mother of $x_i$
5: \hspace{1em} $s \leftarrow D(m)$ \hspace{1em} \text{\texttt{\texttt{\texttt{\texttt{\triangle}}}} Load daughters of $M$
6: \hspace{1em} $\overrightarrow{d} \leftarrow \text{distance}(x_i, s)$
7: \hspace{1em} $J \leftarrow \{j|\overrightarrow{d}[j] < d_c\}$
8: \hspace{1em} if $J \neq \emptyset$ then
9: \hspace{1.5em} $\forall j \in J, \rho^g_j(B^g_j) \leftarrow \rho^g_j(B^g_j) + 1$
10: \hspace{1em} else
11: \hspace{1.5em} $S^g \leftarrow S^g \cup B^g_k(x_i, d_c)$ \hspace{1em} \text{\texttt{\texttt{\texttt{\texttt{\triangle}}}} Create new ball
12: \hspace{1.5em} $\rho^g_k(B^g_k(x_i, d_c)) \leftarrow 1$
13: \hspace{1.5em} $L \leftarrow \{l|\text{distance}(B_k(x_i, d_c), S^{g-1}[l]) < d_c(1 + R)\}$
14: \hspace{1.5em} Update $F^g_{tree}$ \hspace{1em} \text{\texttt{\texttt{\texttt{\texttt{\triangle}}}} Record that $B(x_i, d_c)$ is attached to $\{B_l\}_{l \in L}$
15: \hspace{1.5em} $J \leftarrow k$
16: \hspace{1em} end if
17: \hspace{0.5em} Store $J$ in $F^g_{tree}$
18: end for
19: for all $x_i$ frozen at last iteration do
20: \hspace{0.5em} $m \leftarrow M(x_i)$ \hspace{1em} \text{\texttt{\texttt{\texttt{\texttt{\triangle}}}} Load mother of $x_i$
21: \hspace{0.5em} $s \leftarrow D(m)$ \hspace{1em} \text{\texttt{\texttt{\texttt{\texttt{\triangle}}}} Load daughters of $M$
22: \hspace{0.5em} $\overrightarrow{d} \leftarrow \text{distance}(x_i, s)$
23: \hspace{0.5em} $\forall j \in J, \rho^g_j(B^g_j) \leftarrow \rho^g_j(B^g_j) + 1$
24: end for
25: $TB^g \leftarrow \{B^g_i|\rho^g_i(B^g_i) < N_{min}\}_{i}$
26: return $\{F_{tree}, \{\rho^g_k\}_{k}, S^g, TB^g\}$
27: end procedure
\( R = \frac{3}{4} \) appeared to be a good trade-off according to the conducted experiments.

In order to quickly cover the domain with few balls in the first iteration, \( d_1^c \) can be set to 1. It is justified by the fact that, since the dataset is normalized, \( d_1^c = 1 \) is equal to the standards deviations \( \{\sigma_j\}_{j=1,...,p} \) in each dimension. Moreover the choice \( d_0 \) has little impact then we do not need an accurate normalization. Therefore, we can only pseudo normalize the dataset, that is to say, estimate \( \{\hat{\mu}_j\}_{j=1,...,p} \) then \( \{\hat{\sigma}_j\}_{j=1,...,p} \) with a small subset part of the sample only and compute \( x_{ij} = \frac{x_{ij}}{\hat{\sigma}_j}, \forall (i,j) \in N \times P \).

4 Cluster centers selection and labelisation using \( F_{\text{tree}} \)

This step mainly consists in finding, for each terminal ball, the closest one with higher density. Even if the number of points to be treated is decreased, it can remain too large to be directly processed by the standard peak detection presented in [17]. Instead of computing all the distances between all terminal balls, we use the localisation information provided by \( F_{\text{tree}} \) to compute distances for a few of them only.

This part is based on the following observation. Consider the \( k^{\text{th}} \) ball \( B_{kG}^{-1} \) of generation \( G - 1 \) (where \( G \) is the last generation) and also consider its daughters \( A_k^G = D(B_{kG}^{-1}) \). We define \( C_{kG}^{-1} \) the "champion" of \( B_{kG}^{-1} \), the ball of \( A_k^G \) who has the higher density. Then, excepted for the champion \( C_{kG}^{-1} \), every daughter ball \( B_j^G \in A_k^G \) possess a neighbor of higher density that is also belonging to \( A_k^G \). For these ones, we can restrict the search to those few sister balls. However, for the champion \( C_k^G \), we need to search farther. To do so, we apply quite similar steps between the champions: We first search among the sisters balls of \( B_{kG}^{-1} \) and find \( B_{lG}^{-1} \) (if it exists), the closest one which contains a higher density champion. We then search for the NHPD among the daughter balls of \( B_{lG}^{-1} \). If there isn’t any sister ball containing a higher density champion, we have to search farther again with the same procedure.

The complete algorithm of cluster center selection and labelisation consists in the fol-
lowing steps:

1) Champions election (see Algorithm 5)

This procedure constructs two new objects that share the same structure than $F_{\text{tree}}$. For each ball $B_k^g$, $\rho_{\text{champ},k}^g := \rho_{\text{champ}}^g(B_k^q)$ provides the density of its champion. $c_k^g \in \{0, \ldots, G-1\}$ indicate the champion’s level of $B_k^g$. First, the $c_k^g$ are initialized according to their generation ($c_k^g = G - g$). Then we begin at generation $G - 1$. For each ball $B_k^{G-1}$, we find among its daughters the ball $B_c^G$ which hold the highest density then we set the champion’s level $c_c^g$ of $B_c^G$ to $G - (G - 1) = 1$ and update the champion-densities by setting $\rho_{\text{champ}}^{G-1}(B_k^{G-1}) \leftarrow \rho^G(B_c^G)$. Afterwards, for each ball $B_k^{G-2}$ of the generation $G - 2$, we find among its daughters the ball $B_c^{G-1}$ which hold the highest champion-density then set the champion’s level of the champion carried by $B_c^{G-1}$ to $G - (G - 2) = 2$ and update the champion-densities by setting $\rho_{\text{champ}}^{G-2}(B_k^{G-2}) \leftarrow \rho_{\text{champ}}^{G-1}(B_c^{G-1})$. This procedure is repeated for generations $G - 3, \ldots, 1$.

2) Constructions of $\delta_i$

For each terminal ball, the Tree-Dive procedure (see Algorithm 6) is applied in order to obtain its NHPD and the distance $\delta$ separating them. For a level $c$ champion ball $B$, this algorithm executes the following steps. It first load the ball $B_i^{G-c}$ that has promoted $B$ to the level $c$. Afterwards, the mother $B_k^{G-c-1} = M(B_i^{G-c})$ of $B_i^{G-c}$ and then the daughters $s = D(B_k^{G-c-1})$ of $B_k^{G-c-1}$ are loaded. $B_j^{G-c} \in s$, the closest one with higher champion-density than $\rho(B)$, is retained. There always exist a ball with higher champion’s density, otherwise its champion’s level should be at least $c + 1$. If $B_j^{G-c}$ is a terminal ball then the algorithm stops, if not the daughters of $B_j^{G-c}$ are loaded and the closest one from $B$ with higher champion-density is selected. If it is terminal then it stops, if not it keep diving into the tree until it falls into a terminal ball with higher density. The distance $\delta(B)$ separating this terminal ball and $B$ is finally returned.
3) Cluster center selection and labelisation

\[ \gamma_i^g = \delta_i^g, \rho_i^g \]

are computed, sorted and the turning-point-based decision rule from [11] presented in related work is applied to obtain the cluster centers. The selected centroids take each an unique label and the remaining balls receive the same label than their NHPD. Finally, each data point are labelled as the closest terminal ball containing it.

The complete procedure of IDPC is summarized in Algorithm 7.

Algorithm 5 Champions election

1: procedure ChampElect \((F_{\text{tree}}, \{\rho_k^g\}_g,k, \{B_k^g\}_g,k)\)
2: \hspace{1em} for \(g \in \{G - 1, \ldots, 1\}\) do
3: \hspace{2em} for all \(B_i^g\) do
4: \hspace{3em} if \(B_i^g\) is terminal then
5: \hspace{4em} \(\rho_i^{g+1}(B_i^g) \leftarrow \rho_i^g(B_i^g)\)
6: \hspace{4em} \(c_i^g \leftarrow G - g\) \hspace{2.5em} \(\triangleright\) Upgrade champions level
7: \hspace{2em} else
8: \hspace{3em} \(B_j^{g+1} \leftarrow \text{argmax}_{B_i^{g+1} \in D(B_i^g)}(\rho_i^{g+1}(B_i^{g+1}))\)
9: \hspace{3em} \(\rho_i^{g+1}(B_i^g) \leftarrow \rho_{\text{champ},i}(B_i^{g+1})\)
10: \hspace{3em} \(c_j^{g+1} \leftarrow G - g\)
11: \hspace{2em} end if
12: \hspace{1em} end for
13: \hspace{1em} end for
14: return \((\{\rho_k^{g+1}\}_g,k, \{c_k^g\}_k,g)\)
15: end procedure

5 Experimental results

The generated dataset is composed by 1024508 bidimensional points in the rectangular domain \([-300, 300] \times [-300, 300]\). It contains 13 clusters, its distribution can be seen in the figure 1. The ICMDW has been applied with \(N_{\min} = 0.1\%N\) and \(R = \frac{3}{4}\).

It produced a set of 9224 balls (figure 2), including the 4622 terminal balls (figure 3), all generations confounded. The total number of generations was 16.
Algorithm 6 Tree-dive

procedure TREE-DIVE($B, F_{\text{tree}}, c, \{\rho^g_{\text{champ,k}}\}_{g,k}$)
    $T \leftarrow False$
    Load $B_M$  \hspace{1cm} ▷ mother of the ball that promoted $B$ to level $c$
    while $T == False$ do
        Load $s \leftarrow D(B_M)$  \hspace{1cm} ▷ daughters of $B_M$
        $s_c^+ \leftarrow \{i \mid \rho^c_{\text{champ,i}}(B^c_i) > \rho(B)\}$
        $j \leftarrow \text{argmin}_i(\text{distance}(B, s_c^+[i]))$
        if $B^c_j$ is a terminal Ball then
            $T \leftarrow True$
        else
            $B_M \leftarrow B^c_j$
            $c \leftarrow c + 1$
        end if
    end while
    nhpd $\leftarrow B^c_j$
    $\delta \leftarrow \text{distance}(B, B^c_j)$
    return $\{\delta, \text{nhpd}\}$
end procedure

The density map is represented in the figure 4 over two different points of view. To construct this graphic we have created a grid of points covering the domain and associated, for each point of the grid, the density of the closest terminal ball containing it, to finally plot the surface. The density is well represented however there is some moderately dense balls which are supposed to carry a null or very low density. This problem that we call aliasing occurs in low density areas located near an abrupt peak. An aliasing ball is a ball that has been created lately (at a given generation), preventing it to increase its density count with the points that have been drawn before its creation. Therefore it is set terminal and get back its density at the density reevaluation phase. Even if the density map doesn’t reflect perfectly the true density, the aliasing can only create relatively low density balls (because of the large width of the window $d^0_g$) near a much higher density peak. So its impact on clustering is almost inexistent.

The cluster centers can now be selected. We can see in the decision graph (figure 5) 13 points standing out the others. They are automatically detected with the turning-point
Algorithm 7 IDPC

procedure IDPC
    \[ N_{\text{min}} \leftarrow 0.1\%N \]
    \[ d^1_c \leftarrow 1 \]
    Pseudo Normalize
    Randomize
    \[
    \begin{align*}
    &\{\{B^g_k\}_{g,k}, F_{\text{tree}}, \{\rho^g_k\}_{g,k}, \{TB^g_k\}_g, \{cc_i\}_i\} \leftarrow ICMDW(d^1_c, N_{\text{min}}, R) \\
    &\{\{\rho_{\text{champ},k}^g\}_{g,k}, \{c^g_k\}_{k,g}\} \leftarrow \text{CHAMPELECT}(F_{\text{tree}}, \{B^g_k\}_{g,k}, \{\rho^g_k\}_{g,k}) \\
    \end{align*}
    \]
    for all \((g, k)\) do
        \[
        \{\delta^g_k, \text{nhpd}_{g,k}\} \leftarrow \text{TREE-DIVE}(B^g_k, F_{\text{tree}}, c^g_k, \{\rho_{\text{champ},k}^g\}_{g,k})
        \]
    end for
    \[ \gamma \leftarrow \text{sort}(\delta, \rho) \]
    for \(i \leftarrow \{1, \ldots, m\}\) do
        \[ k^1_i \leftarrow \frac{\gamma_i - \gamma_1}{\gamma_i - \gamma_{i-1}} \]
        \[ k^{i-1}_i \leftarrow \frac{\gamma_{i+1} - \gamma_i}{1} \]
    end for
    \[ tp \leftarrow \arg\max_i \{k^{i-1}_i\} \quad \triangleright \text{Turning-point} \]
    for all \(j \leq tp\) do
        Label\([j]\) \leftarrow \(j\)
    end for
    for all \(j \geq tp + 1\) do
        Label\([j]\) \leftarrow Label[\text{nhpd}_j] \quad \triangleright \text{Labelise Balls} \]
    end for
    for \(i \leftarrow \{1, \ldots, N\}\) do
        Label\([i]\) \leftarrow Label[cc_i] \quad \triangleright \text{Labelise datapoints} \]
    end for
    return Label
end procedure

procedure. The sorted gamma values are represented in the figure 5 and the selected cluster centers are represented in red and their location and the corresponding clustering results obtained by propagation of the label are showed in the figure 6.

Note that we didn’t have generated any flat clusters (like an uniform) because the original algorithm, and so our, isn’t able to deal with. In those situations, the algorithm may detect several peaks for only one real cluster. In a future work, we will benefit from the reduced set of pretender centroids and their localisation information provided by our algorithm to develop a better cluster center selection. To do so we might not only take into account the distance separating two peaks but rather the density connectivity between them. In other
words, searching if there is a dense trajectory to reach one peak from another one.
Figure 2. All the balls
Figure 3. Terminal balls only
Figure 4. Density map
Figure 5. Center selection

Figure 6. Clustering results
6 Conclusion and future work

Many extensions of FDPC have been developed but none is able to deal with multidimensional and large datasets that arise in many practical data analysis applications. In this paper we have presented an efficient algorithm able to manage those kind of data.

For that purpose, we have designed an iterative procedure which creates both localisation and density map which gives, in a computationally efficient way, the densities $\rho_i$ and the distances $\delta_i$.

It is adaptive, the results being barely influenced by the few free parameters to be tuned, so that the automatic choice is often satisfactory. This last point is a dramatic improvement over most of the existing algorithms that tend to depend critically on the initial conditions.

An important remaining work to be performed is the recoding of the procedure in a compiled language with optimized data structures, from which a gain of at least an order of magnitude is expected. Moreover we aim to develop a better cluster selection unlocked by our intermediate results, able to manage flat distributions. Finally, we expect to delete the aliasing in the density map.

References


[22] Yi Wang, Qixin Chen, Chongqing Kang, and Qing Xia. Clustering of electricity consumption behavior dynamics toward big data applications.


