Supplementary material for: Data with Density-Based Clusters: A Generator for Systematic Evaluation of Clustering Algorithms

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The supplement expands on the related work included in "Data with Density-Based Clusters: A Generator for Systematic Evaluation of Clustering Algorithms" in Appendix A. This includes a more in-depth look at the other data generators in Appendix A.1 and a detailed description of the clustering algorithms used in Appendix A.2.

The supplement also gives additional explanations on the *equivalence* of datasets in Appendix B.

Furthermore, we include results and background for an extended benchmark study with 6 additional algorithms and an additional basic data setting in Appendix C. All datasets, including the 2d visualization dataset, as well as the experiment logs for this evaluation, can be found in the GitHub repository of the paper: https://github.com/PhilJahn/DENSIRED.

A Expanded Related Work

A.1 Data Generators

There are data generators for many different settings [1], e.g., for multivariate ordinal data [2], for subspace clustering [3], and for text, trees, and logical interpretations [4]. Thus, data-generating functions have been established as a useful tool for a controlled evaluation of procedures in a plethora of domains ranging from healthcare [5, 6], routing problems [7], flow networks [8], to manufacturing [9].

In [10], the authors use a "self-implemented data generator" as the basis for their experiments for evaluating a novel density-based subspace clustering. However, neither implementation nor exact specifics that are sufficient to reproduce the datasets are given.

Pei's data generator described in [11] offers the ability to create clusters of different shapes based on difficulty levels. This includes clusters that could be suitable for evaluating density-based clustering algorithms. However, as the approach is

limited to two-dimensional data, it can not be used to evaluate a method's quality in higher dimensional space.

Milligan's method introduced in [12,13] sets up cluster boundaries and fills them based on a random distribution around the midpoints of the boundary lengths. This approach was augmented in [14], adding the ability to specify the degree of separation and allowing arbitrary positive definite matrices as covariance matrices. Nonetheless, the data is limited to what can be generated by a distribution around a singular midpoint.

MixSim [15] is a popular data generator for overlapping clusters. It simulates Gaussian mixture models with varying covariance matrices. Using an inverse Box-Cox transformation, MixSim can also provide non-normally distributed data. However, MixSim only uses a single Gaussian distribution per component and clusters' shapes are limited by the features of inverse Box-Cox transformations. This, in turn, limits MixSim to non-arbitrary shapes. Still, MixSim has been used to evaluate density-based clustering in [16].

Scikit-learn [17] offers several ways to generate data, though most aim towards 2-dimensional data. Primarily of interest are the blobs and swiss-roll generators, which have been used to evaluate density connectivity-based clustering methods [18]. The blobs generator produces isotropic Gaussian distributions that are either randomly placed or have their center specified by user input. However, the generator is limited to purely Gaussians, and can thus not create arbitrarily shaped clusters. The swiss-roll generator is limited to 3 dimensions and does not generate separate ground-truth clusters but rather a single distribution that is shaped like a roll.

Clugen [19] uses line segments as a basis for generating clusters with directionality. It builds on top of a 2-dimensional data generator [20] but expands the method to a multi-dimensional setting. Clugen is based on arbitrary statistical distributions. As clusters have only a single supporting line, their overall shape can not be arbitrarily shaped.

OCLUS [21] offers a data generation approach for overlapping clusters, expressed by shared density between clusters. The clusters are based on user-defined random distributions, but as these distributions only apply independently in distinct dimensions, clusters are constrained to be of non-arbitrary shape.

HAWKS [22] is a data generation method based on an evolutionary algorithm. HAWKS creates Gaussian clusters where the distributions are adjusted based on mutation and cross-over from parent datasets. Constraints can be applied that are used to select these parents, in order to target specific properties for the final dataset. Still, Gaussian clusters can not be arbitrarily shaped.

MDCGEN [23] is a data generator for subspace clusters. It uses hyper-grids for overlap control and cluster positioning and can produce radial-based or multivariate clusters. The coordinates of a cluster's points are drawn from a distribution for each dimension. This includes ring-shaped distributions that can be considered density-connected. However, this leads to only a very limited range of density-connected clusters, which is not sufficient for a broad systematic evaluation of density-based clustering algorithms: the crucial benefits of these cluster-

ing methods can primarily be seen and evaluated on arbitrarily shaped clusters, which can not be generated with MDCGEN.

SynDECA [24] offers to create clusters of different shapes. Before creating the actual clusters, bounding boxes are determined to prevent any overlap. SynDECA randomly chooses a shape type for each bounding box. Here, the irregular shapes are especially interesting. SynDECA offers two ways of generating irregular shapes. Irregular 1 performs a type of random walk that expands a random data point with an ε -range by further points with the same radius. This could, however, violate the bounding box and overlap with other clusters. Furthermore, this approach offers very little control. Irregular 2 works by splitting the bounding box into smaller boxes. Starting from a random box, an adjacent box is selected and randomly filled with points. This continues until a threshold of data points is reached. While this produces complex-shaped clusters, the usage of hyperrectangular bounding boxes makes the clusters easily separable despite the actual shape.

DataGen [25] is a data generator that first generates class boundaries based on an initial random assignment of data points which serve as training data for a classifier. The decision boundaries of this classifier are used to define class regions which are then populated with data points following a random distribution. While powerful, DataGen is limited by the decision boundaries of the classifier; there are no guarantees that regions are continuously dense or even connected. Furthermore, users can not control most properties of the data.

Seed Spreader [26, 27] was already used [26, 28] to create higher-dimensional density-connected clusters. It works by generating data points with a fixed radius (based on either a global radius value or on the current cluster number [27]) surrounding the current position of the random walk. It either samples until a predefined amount of points within the hypersphere is reached or - based on a preset chance - jumps to a random position in the data space, at which the hypersphere sampling is terminated early. If the full amount of points has been generated from the hypersphere, the random walk moves on. Seed Spreader has multiple issues that DENSIRED addresses (i.e., a lack of user control, potentially overlapping clusters, and no knowledge about the number of clusters prior to the generation), which are covered in the main paper.

Availability Availability of a generator is a very important aspect for reproducible evaluation and benchmarking.

Fachada and de Andrade [19] already noted this importance in their paper on Clugen. They made Clugen available in multiple programming languages⁵ and also documented the availability of related data generators.

Pei's generator [11] is, to the best of our knowledge, not publicly available [19]. An implementation of Milligan's method [12] is available as a C++ implementation [19, 29]⁶, but the implementation is not usable out of the box [19].

The improved version of Milligan's method [14] is available as an R implemen-

⁵ https://github.com/clugen, last accessed: Mar 22nd, 2024

⁶ https://clusutils.sourceforge.net/, last accessed: Mar 22nd, 2024

tation $[19]^7$.

MixSim [15] is also available as an R implementation [15, 19]⁸.

The Scikit-learn library is public, and as such both the blobs as well as the swiss-roll generators are easily accessible in Python⁹.

OCLUS [21] is, to the best of our knowledge, not publicly available, though its code can be requested from the authors [19].

A Python implementation of HAWKS [22] is available $[19, 22]^{10}$.

MDCGen is available in both Python and MATLAB [19,23]¹¹.

SynDECA appears to have been available at some point [19]¹², but the link is no longer functional, making the method inaccessible.

The authors of DataGen [25] expressed their intention of making their code available in the UCI repository [30], but we have not been able to find the generator. To the best of our knowledge, Seed Spreader was not made publicly available alongside the publications describing it [26, 27]. However, our Python reimplementation based on the provided description can be found in our repository. Our method, DENSIRED, is available there as a Python implementation as well.

A.2 Clustering Methods

We introduce the clustering methods used in the expanded benchmarking in Appendix C in the following.

Centroid-based Clustering k-Means [31] is one of the most fundamental clustering algorithms and requires only the number of clusters as user input. Its goal is to minimize the distances of all points to their respective cluster centroids, which is most often done in an iterative, non-deterministic way.

LDA-Km [32] is an extension of k-Means that incorporates linear discriminant analysis (LDA) [33]. LDA is used to select a suitable subspace for clustering, while the clustering itself is performed by k-Means. As both methods optimize the same objective function, subspace and cluster adjustments are performed simultaneously.

X-Means [34,35] is an adaption of k-Means that integrates information-theoretic criteria to maximize the model's quality. The major advantage besides its efficiency is that X-Means finds the number of clusters automatically.

 $^{^7}$ https://cran.r-project.org/web/packages/clusterGeneration/index.html, last accessed: Mar 22nd, 2024

 $^{^8}$ https://cran.r-project.org/web/packages/MixSim/index.html, last accessed: Mar 22nd, 2024

https://scikit-learn.org/stable/modules/generated/sklearn.datasets. make_blobs.html and https://scikit-learn.org/stable/modules/generated/sklearn.datasets.make_swiss_roll.html, last accessed: Mar 22nd, 2024

 $^{^{10}}$ https://github.com/sea-shunned/hawks, last accessed: Mar 22nd, $2024\,$

https://github.com/CN-TU/mdcgenpy and https://github.com/CN-TU/mdcgen-matlab, last accessed: Mar 22nd, 2024

 $^{^{12}}$ https://sites.google.com/site/syndeca/, last attempted access: Mar 22nd, 2024

Spectral Clustering

Spectral Clustering [36–38] is another fundamental clustering concept. It is based on the eigendecomposition of a similarity graph representing the data. Points are then clustered in their spectral embedding, where often k-Means is used. SCAR [39] is a novel spectral clustering method working on weighted kNN graphs. It removes noisy edges from the affinity graph building on top of [40] while accelerating the eigendecomposition with the Nyström method [41]. SpectACl [42] combines spectral and density-based clustering by maximizing the average cluster density. It is based on the ratio cut of the normalized ϵ -neighborhood graph given by its spectrum.

Hierarchical Clustering Agglomerative clustering methods like [43–46] are bottom-up hierarchical clustering methods that are based solely on some similarity measure between points. They differ mainly in the way they measure similarity between groups of points, where single-linkage, average-linkage, complete-linkage, and Ward's method [43] are the most common methods. Ward was the method chosen for the evaluation in Appendix C (as well as for the main paper). Birch [47] is a hierarchical clustering approach that is based on a height-balanced tree with so-called *Clustering Features* as leaves. Areas of higher density are considered (nested) subclusters, while points in sparse areas are removed as outliers. Clustering features contain information on the number of data points as well as the linear and squared sums of their values. Each non-leaf node corresponds to a higher-level cluster containing its children.

Deep Clustering

In recent years, deep clustering approaches that combine a clustering objective with neural networks have become increasingly popular. DEC [48] works by simultaneously learning a feature representation of the data and their cluster assignments using the Kullback-Leibler divergence. It is initialized as a stacked autoencoder, but only the encoder part is retained to produce the feature embedding for clustering.

IDEC [49] is an extension of DEC that keeps the decoder part of the autoencoder to retain the local structure and to ensure that the representation is not distorted. The clustering loss that is used to obtain an embedding conductive of clustering is directly applied to the embedding, whereas the decoder is untouched and has to be able to reconstruct the data again.

Unlike DEC and IDEC, DipDECK [50] is capable of estimating the number of clusters within the latent space of an autoencoder. It does so by using Hartigans' dip test [51] after heavily overestimating the number of clusters.

The DipEncoder [52] uses a gradient formulation of Hartigans' dip values [51] to separate multimodal data in the embedded space. The cluster assignment is achieved on the so-called modal interval, which defines the main mode of the data.

Density-based Clustering

Mean Shift [53] aims at finding modes of an underlying density function by recursively applying the eponymous mean shift procedure [54,55]. A cluster contains the data points visited by all the mean shift procedures after they converged to the same location. Data points visited by multiple procedures are assigned to the predominant one.

The Density Peak Clustering Algorithm (DPCA) [56] finds clusters that are centered around a local density maximum by regarding the distances of points to the closest point of higher density. Cluster assignments are based on the cluster of the nearest neighbor with higher density.

DCF [57] uses the concept of modal sets introduced in [58]. It determines cluster cores for instances with the highest densities based on kNN with similar densities. This leads to separated cluster cores that imply the clusters.

Data Density Based Clustering (DDC) [59] uses the highest density points as cluster centers and assigns data points within a cluster radius. Clusters are refined and already assigned samples are removed before repeating the process until the number of considered points falls below a threshold.

Density-Connectivity-based Clustering

DBSCAN [26,60,61] finds density-connected clusters of arbitrary shape by connecting *core* points (i.e., points that have at least minPts neighbors in their ε -range) in a manner similar to Prim's algorithm. It finds the number of clusters automatically and considers noise points.

OPTICS [62, 63] is based on the concepts of DBSCAN and computes an augmented cluster ordering usually visualized as OPTICS plot. This plot allows users to easily find a suitable ε for DBSCAN as well as a DBSCAN-like clustering [62] and gain insights into the cluster structure. In our experiments in Appendix C, OPTICS clusters were automatically obtained from OPTICS plot based on the ξ -steepness [62].

HDBSCAN [64,65] is an extension of DBSCAN that makes use of concepts from hierarchical clustering. The core idea of HDBSCAN is to identify clusters that persist over multiple ε -values, i.e., multiple density levels. The hierarchy of the underlying tree structure of HDBSCAN is determined by the clusters' stability over varying ε -values as well as a minimal cluster size. HDBSCAN aims at optimizing the stability of all clusters while maintaining their disjointness.

Multi-density DBSCAN (MDD) [66] extends DBSCAN to deal with clusters of different densities. It iteratively processes data points with decreasing density and forms clusters by expanding to the k-neighbors with similar densities.

B Expanded Theory, and Concept

B.1 "Density-connectivity"

Our generator creates datasets containing density-connected clusters. We follow the well-known concept from clustering algorithms like DBSCAN [60] or

HDBSCAN [64]. Note that this differs from mode-seeking density-based concepts as, e.g., used in Mean Shift [53]. For completeness, we provide definitions for density-connectivity based on [60,67].

Definition 1. ε -neighborhood and core-property

The ε -neighborhood of a point p in a dataset X is defined as $N_{\varepsilon}(p) = \{q \in X : dist(p,q) \leq \varepsilon\}$ for $\varepsilon > 0$ and a distance measure $dist(\cdot,\cdot)$. A point is a core point w.r.t. $\varepsilon \in \mathbb{R}^+$ and $\mu \in \mathbb{N}$ iff it has at least μ points in its ε -neighborhood.

Definition 2. (Directly) Density-Reachable

A point q is directly density-reachable from a point p for some $\varepsilon > 0$, $\mu \in \mathbb{N}$ iff p is in the ε -neighborhood of q and q is a core point.

A point q is density-reachable from a point p iff there is a chain of directly density-connected core points between them.

Definition 3. Density-Connected

A point p is density-connected to a point q for $\varepsilon > 0$, $\mu \in \mathbb{N}$ iff there is a point o such that p and q are both density-reachable from o.

Definition 4. Density-based Cluster

A (density-based) cluster $C^{\varepsilon,\mu} \subseteq X$ is a "maximal set of density-connected objects" [67].

B.2 Equivalence of datasets

For analyzing centroid-based clustering algorithms, clusters are usually generated by drawing from a Gaussian distribution with predefined centers and variances. The resulting datasets are similar, but not equal. With that intuition, we call such clusters equivalent and define it accordingly for density-based clusterings:

Definition 5. Equivalence of (density-based) clusterings.

Two density-based clusterings \mathcal{A}, \mathcal{B} are equivalent if there is a bijective mapping between them. I.e., there is a bijective mapping $f_c : \mathcal{A} \to \mathcal{B}$ of the clusters such that for every cluster in \mathcal{A} , there is also a bijective mapping of all its points to the points in the associated cluster in $\mathcal{B}: \forall c^{\mathcal{A}} \in \mathcal{A} \exists c^{\mathcal{B}}: f_p : c^{\mathcal{A}} \to c^{\mathcal{B}}$.

Definition 6. Equivalence of density-based datasets.

Two datasets $X \in \mathbb{R}^{n \times d}$, $Y \in \mathbb{R}^{n \times d}$ are equivalent regarding their density-connectivity if, for every ε and μ , there exist δ, ν , such that the ε, μ - clustering of X is equivalent to the δ, ν -clustering of Y.

For an unbiased evaluation of a novel method, it is important to eliminate random effects. Especially for datasets with density-based clusters, these effects can be decisive for the perceived quality of, e.g., a clustering model, see [68]. Using several equivalent datasets allows the elimination of these random effects for a proper analysis and evaluation of novel clustering algorithms and other methods working on similar data.

B.3 Challenge: Generating equivalent clusters

Generating equivalent density-based clusters is a challenging problem. In contrast to centroid-based clustering concepts, density-based clustering has a discrete, non-continuous aspect expressed by the parameter $\mu \in \mathbb{N}$. Thus, changing a single point in a dataset can lead to a drastically different clustering. Especially, clusters that fulfill the density-criterion only tightly can disintegrate into several clusters if a decisive point is missing or changing positions as shown in Figure 1.

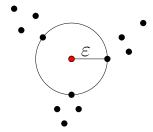


Fig. 1: Consider this dataset and its clustering for $\mu=4$ and ε as given by the radius of the circle. All data points form one cluster. If the central red data point is moved to any other position, the cluster splits up. Without the red data point, there are three density-based clusters in the dataset.

C Expanded Benchmarking

C.1 Experiment Setup

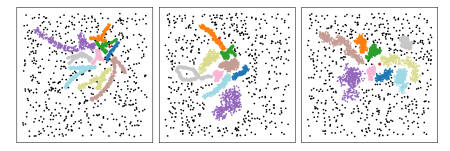


Fig. 2: Two-dimensional benchmark datasets (the cluster shape is more 'stretched' and the right is 'compact'; the middle 'mix' data was only used for the two-dimensional case).

For the expanded benchmark study, we evaluated 20 clustering algorithms on data produced by DENSIRED. For an in-depth analysis of higher-dimensional space, we evaluate the models on two different basic settings with a shared set of parameters besides the dimensionality:

- i) The first category ("stretched") contains branching data with high momentum and larger step size between consecutive cores¹³, yielding more spread out clusters with tendentially higher intrinsic dimensionalities. This setting was also the one employed in the main paper.
- ii) The second category ("compact") contains non-branching data without momentum resulting in compact cluster structures ¹⁴.

The two-dimensional versions for both settings are shown in Figure 2 on the left and right sides. Figure 3 features the corresponding cluster skeletons for the two-dimensional case. For both settings, we generate datasets of varying dimensionality $d \in [2, 5, 10, 50, 100]$ scaled to a range between 0 and 1 by their respective maximal value with 5000 data points for each dataset.

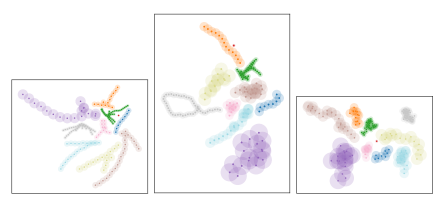


Fig. 3: Skeletons of the two-dimensional benchmark datasets (the cluster shape is more 'stretched' and the right is 'compact'; the middle 'mix' data was only used for the two-dimensional case).

Note that the visualized data represents just a small subset of the generator's capabilities. Aside from the data generated for the "stretched" and "compact" settings, there is also a third 2D dataset "mix", which features a more varied setting of the parameters but was primarily intended for visualization purposes. It is displayed in the middle for both Figure 2 and Figure 3. The datasets used in our experiments are provided as novel benchmark datasets in our repository. We conducted extensive parameter grid searches to find the best possible quality for each model for a fair comparison. In cases where an algorithm requires an ex-

 $^{^{13}}$ Exact values: $\omega=0.8,\,\delta=1.5,\,\beta=0.1,\,\varkappa=1,\,200$ cores

 $^{^{14}}$ Exact values: $\omega=0,\,\delta=1,\,\beta=0,\,\varkappa=0,\,200$ cores

plicit number of clusters, the number of clusters was set to 10 in correspondence to the number of synthesized (density-based) ground-truth clusters. For simplicity, the results of the best-performing setting for each algorithm were chosen and are used here (best mean performance for both "compact" and "stretched", highest clustering performance for "mix"). Best performing was defined as the highest sum of ARI [69] and NMI [70]. A full log of the results for the evaluated parameters in the initial parameter sweep is listed in our repository.

The implementations for k-Means, Mean Shift, Spectral Clustering, Agglomerative Clustering, BIRCH, DBSCAN and OPTICS stem from scikit-learn¹⁵ [17]. The implementations for LDA-Km, X-Means, DEC, IDEC, DipDeck, DipEncoder and MDD originate from ClustPy¹⁶ [71]. The implementation of HDB-SCAN stems from the hdbscan python package¹⁷. DCF¹⁸, SCAR¹⁹, SpectACl²⁰, DPCA²¹, and DDC²² originate from the linked repositories.

C.2 "Stretched"-Setting

The results for the "stretched"-setting for the data generator are listed in Table 1. Figure 4 shows the general tendencies in performance of the algorithms across all examined dimensionalities. The best-performing parameters are described in Table 2.

 $^{^{15}}$ https://scikit-learn.org, last accessed: Oct 5th, 2023

https://github.com/collinleiber/ClustPy/tree/main, last accessed: November 15th, 2023; LDA-Km, X-Means and MDD used the initial v0.0.1-alpha version; DEC, IDEC, DipDeck and DipEncoder used v0.0.2-beta

 $^{^{17}}$ https://github.com/scikit-learn-contrib/hdbscan/tree/master, last accessed: October 5th, 2023

¹⁸ https://github.com/tobinjo96/DCFcluster, last accessed: October 5th, 2023

¹⁹ https://github.com/SpectralClusteringAcceleratedRobust/SCAR, last accessed: November 4th, 2023

²⁰ https://bitbucket.org/Sibylse/spectacl/src/master/, last accessed: October 11th, 2023

²¹ https://github.com/colinwke/dpca, last accessed: October 31st, 2023

²² https://github.com/RHyde67/DDC-Python, last accessed: November 16th, 2023

Setting																				
dimensionality		2-d			5-dim			10-dim			50-dim			100-dim						
Metric		ARI	N	NMI	1	ARI	1	NMI		ARI		NMI		ARI	1	NMI	1	ARI	l N	IMI
k-Means		± 0.04	0.62	± 0.02	0.32	± 0.02		± 0.01				±0.03		±0.04		±0.02		± 0.05		
LDA-Km	0.48	± 0.03	0.61	± 0.02	0.31	± 0.01		± 0.01	0.60	± 0.06		± 0.04		± 0.03		±0.02	0.67	± 0.00	0.80	± 0.00
X-Means		± 0.02	0.63		0.35	± 0.02		± 0.02		± 0.03		± 0.02		± 0.01		±0.01		± 0.05		± 0.02
Spectral Clust.	0.57			± 0.00	0.43	± 0.00		± 0.00		± 0.00		±0.00		± 0.02		± 0.01		± 0.00		± 0.00
SCAR		± 0.02		± 0.01	0.47			± 0.02		± 0.00		± 0.00		± 0.02		± 0.02		± 0.00		± 0.00
SpectACl		± 0.04		± 0.02	0.54	± 0.04	0.80	± 0.02		± 0.02		± 0.01		± 0.02	0.87			± 0.04		± 0.02
Ward Clust.	0.43		0.59		0.35		0.57		0.61		0.75		0.66		0.80		0.72		0.85	
Birch	0.49		0.61		0.27		0.49		0.61		0.74		0.66		0.80		0.72		0.85	
DEC	0.36	± 0.06	0.51	± 0.05	0.41	± 0.02	0.61		0.72	± 0.05	0.85	± 0.03	0.59		0.77	± 0.04	0.52	± 0.06	0.75	± 0.05
IDEC	0.43		0.55	± 0.05	0.40	± 0.04	0.60	± 0.03	0.72		0.85	± 0.03	0.64		0.80		0.73	± 0.06	0.83	± 0.03
DipDECK	0.52	± 0.06	0.62		0.34	± 0.03	0.52			± 0.06	0.76			± 0.05		±0.03	0.77			± 0.04
DipEncoder	0.42	± 0.04	0.60	± 0.02	0.42	± 0.05	0.57	± 0.02	0.73	± 0.03	0.84	± 0.02	0.66	± 0.05	0.79	±0.02	0.69	± 0.10	0.83	± 0.06
Mean Shift	0.64		0.68		0.44		0.62		0.73		0.76	,	0.83	5	0.80	<mark>)</mark>	0.87		0.82	
DPCA	0.43		0.58		0.19		0.41		0.61		0.72		0.77	<u> </u>	0.80)	0.82		0.85	
DCF	0.91		0.90		1.00		0.99		1.00		0.99		0.95	,	0.88	3	0.95		0.88	
DDC	0.42		0.64		0.34		0.63		0.63		0.72		0.65	,	0.69)	0.78		0.69	
DBSCAN	0.87		0.86		0.93		0.91		0.92		0.94		1.00)	1.00)	1.00		1.00	
OPTICS	0.89		0.93		0.91		0.95		1.00		1.00		1.00)	1.00	<u> </u>	1.00		1.00	
HDBSCAN	0.81		0.80		0.95		0.94		1.00		1.00		1.00)	1.00)	1.00		1.00	
MD-DBSCAN	0.96		0.95		1.00		1.00		1.00		1.00		1.00	,	1.00	<u> </u>	1.00		1.00	
	NMI											Al	RI							
0.8 0.6 0.8 0.8 0.8 0.8 0.8 0.8 0.8 0.8 0.8 0.8						•	0.8- 0.6- 0.4- 0.2-	Ī		Ī					₽ .					

Table 1: Clustering results for different dimensionalities for the "stretched"-Setting

Fig. 4: NMI (left) and ARI (right) across all examined dimensionalities for the setting from the "stretched" setting. The coloration indicates the algorithm category as described in Appendix A.2.

C.3 "Compact"-Setting

The results for the "stretched"-setting for the data generator are listed in Table 3. Figure 5 shows the general tendencies in performance of the algorithms across all examined dimensionalities. The best-performing parameters are described in Table 4.

C.4 "Mix"-Setting

The "mix"-setting did not include any higher-dimensional datasets. It was primarily intended for visualization purposes. The cluster assignments for the best-performing clustering runs are displayed in Figure 6, where best-performing means the highest sum of NMI and ARI. The best-performing parameters are described in Table 5.

Table 2: Best performing parameters (out of the evaluated ones) for different dimensionalities for the "stretched"-Setting

dimensionality	2-dim	5-dim	10-dim	50-dim	100-dim
c-Means	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,
c-ivicans	n_init: 2	n_init: 2	n_init: 3	n_init: 7	n_init: 6
DA-Km	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,
JDA-KIII	n_init: 1	n_init: 1	n_init: 1	n_init: 2	n_init: 6
	n_clusters_init: 8,	n_clusters_init: 6,	n_clusters_init: 7,	n_clusters_init: 3,	n_clusters_init: 2,
X-Means	max_n_clusters: 10,	max_n_clusters: 28,	max_n_clusters: 10,	max_n_clusters: 10,	max_n_clusters: 17,
	n split trials: 3	n split trials: 8	n split trials: 1	n split trials: 2	n split trials: 10
	n clusters: 10,	n clusters: 10,	n clusters: 10,	n clusters: 10,	n clusters: 10,
Spectral Clust.	affinity: 'nearest neighbors',	affinity: 'nearest neighbors',	affinity: 'nearest neighbors',	affinity: 'nearest neighbors',	affinity: 'nearest neighbor
•	nearest neighbors: 25	nearest neighbors: 75	nearest neighbors: 120	nearest neighbors: 140	nearest neighbors: 95
	k: 10,	k: 10,	k: 10,	k: 10.	k: 10,
	nn: 15,	nn: 15,	nn: 100,	nn: 100,	nn: 50,
	alpha: 0.9,	alpha: 0.9,	alpha: 0.9,	alpha: 0.7,	alpha: 0.8,
CAR	theta: 1000	theta: 2000	theta: 2000	theta: 3000	theta: 2000
	weighted: True,	weighted: True,	weighted: True,	weighted: True,	weighted: True,
	laplacian: 0,	laplacian: 0,	laplacian: 0,	laplacian: 0,	laplacian: 0,
	normalize: True	normalize: True	normalize: True	normalize: True	normalize: True
	n clusters: 10,	n clusters: 10,	n clusters: 10,	n clusters: 10,	n clusters: 10,
SpectACl	affinity: 'radius neighbors',	affinity: 'radius neighbors',	affinity: 'radius neighbors',	affinity: 'radius neighbors',	affinity: 'radius neighbors
pectaci	epsilon: 0.12	epsilon: 0.04	epsilon: 0.1	epsilon: 0.2	epsilon: 0.2
		n clusters: 10,	n clusters: 10,		
Ward Clust.	n_clusters: 10,			n_clusters: 10,	n_clusters: 10,
	linkage: 'ward'	linkage: 'ward'	linkage: 'ward'	linkage: 'ward'	linkage: 'ward'
	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,
Birch	threshold: 0.05,	threshold: 0.01,	threshold: 0.01,	threshold: 0.01,	threshold: 0.01,
	branching_factor: 75	branching_factor: 150	branching_factor: 2	branching_factor: 2	branching_factor: 2
	n_clusters : 10,	n_clusters : 10,	n_clusters : 10,	n_clusters : 10,	n_clusters : 10,
DEC	alpha: 0.1,	alpha: 0.1,	alpha: 0.1,	alpha: 0.1,	alpha: 0.1,
	embedding_size: 2	embedding_size: 5	embedding_size: 10	embedding_size: 2	embedding_size: 2
	n_clusters : 10,	n_clusters : 10,	n_clusters : 10,	n_clusters : 10,	n_clusters : 10,
IDEC	cluster_loss_weight: 0.01,	cluster_loss_weight: 1.0,	cluster_loss_weight: 1.0,	cluster_loss_weight: 1.0,	cluster_loss_weight: 1.0,
DEC	alpha: 0.1,	alpha: 0.1,	alpha: 0.1,	alpha: 0.2,	alpha: 1.0,
	embedding_size: 2	embedding_size: 5	embedding_size: 10	embedding_size: 2	embedding_size: 2
	cluster loss weight: 0.01,	cluster loss weight: 0.01,	cluster loss weight: 0.01,	cluster loss weight: 0.1,	cluster loss weight: 0.1,
	dip merge threshold: 0.95,	dip merge threshold: 0.95,	dip merge threshold: 0.75,	dip merge threshold: 0.95,	dip merge threshold: 0.7
DipDECK	embedding size: 2,	embedding size: 2,	embedding size: 10,	embedding size: 2,	embedding size: 2,
•	n clusters init: 10,	n clusters init: 15,	n clusters init: 10,	n clusters init: 15,	n clusters init: 15,
	max n clusters: 15	max n clusters: 20	max n clusters: 15	max n clusters: 20	max n clusters: 20
	n clusters: 10,	n clusters: 10,	n clusters: 10,	n clusters: 10,	n clusters: 10,
DipEncoder	rec. loss weight: 10.0,	rec. loss weight: 100.0,	rec. loss weight: 10.0,	rec. loss weight: 100.0,	rec. loss weight: 10.0,
31p12meoder	embedding size: 2	embedding size: 5	embedding size: 10	embedding size: 2	embedding size: 2
	cluster all: True,	cluster all: True,	cluster all: True,	cluster all: True,	cluster all: True,
Mean Shift	bandwidth: 0.06	bandwidth: 0.06	bandwidth: 0.15	bandwidth: 0.17	bandwidth: 0.19
	density threshold: 0.01,	density threshold: 0.01,	density threshold: 0.01,	density threshold: 0.01,	density threshold: 0.01,
DPCA	distance threshold: 0.05,	distance threshold: 0.15,	distance threshold: 0.2,	distance threshold: 0.2,	distance threshold: 0.2,
JI CA					
	gauss_cutoff: True	gauss_cutoff: True	gauss_cutoff: True	gauss_cutoff: False	gauss_cutoff: True
OCF	k: 7,	k: 10,	k: 20,	k: 40,	k: 40,
	beta: 0.991	beta: 0.96	beta: 0.98	beta: 0.99999	beta: 0.99999
DDC	radius: 0.06,	radius: 0.08,	radius: 0.2,	radius: 0.22,	radius: 0.24,
	merge: True	merge: True	merge: True	merge: True	merge: True
DBSCAN	epsilon: 0.008,	epsilon: 0.015,	epsilon: 0.03,	epsilon: 0.09,	epsilon: 0.09,
	min_samples: 3	min_samples: 2	min_samples: 4	min_samples: 2	min_samples: 2
	cluster_method: 'xi',	cluster_method: 'xi',	cluster_method: 'xi',	cluster_method: 'xi',	cluster_method: 'xi',
DTICE	xi: 0.05,	xi: 0.04,	xi: 0.15,	xi: 0.08,	xi: 0.3,
OPTICS	min_cluster_size: 200,	min_cluster_size: 150,	min_cluster_size: 50,	min_cluster_size: 100,	min_cluster_size: 10,
	predecessor_corr.: True	predecessor_corr.: True	predecessor_corr.: True	predecessor_corr.: True	predecessor_corr.: True
IDBSCAN	min cluster size: 6	min cluster size: 5	min cluster size: 5	min cluster size: 6	min cluster size: 2
	var: 2.5,	var: 1.8,	var: 1.5,	var: 1.5,	var: 1.2,
MD-DBSCAN	k: 6,	k: 10,	k: 15,	k: 40,	k: 40,

Table 3: Clustering results for different dimensionalities for the "compact"-Setting

dimensionality	2-dim		5-dim		10-dim		50-0	lim	100-dim	
Metric	ARI	NMI								
k-Means	0.77 ± 0.01	0.78 ± 0.01	0.73 ± 0.05	0.80 ± 0.02	0.72 ± 0.01	0.80 ± 0.01	0.76 ± 0.04	0.85 ± 0.02	0.66 ±0.06	0.83 ± 0.04
LDA-Km	0.78 ± 0.00	0.79 ± 0.00	0.72 ± 0.00	0.81 ± 0.00	0.70 ± 0.00	0.80 ± 0.00	0.74 ± 0.00	0.84 ± 0.00	0.63 ±0.00	0.81 ± 0.00
X-Means	0.84 ± 0.00	0.83 ± 0.00	0.86 ± 0.03	0.86 ± 0.01	0.81 ± 0.04	0.85 ± 0.01	0.94 ± 0.04	0.96 ± 0.02	0.90 ±0.06	0.95 ± 0.03
Spectral Clust.	0.78 ± 0.00	0.84 ± 0.00	0.67 ± 0.00	0.85 ± 0.00	0.65 ± 0.00	0.80 ± 0.00	0.77 ± 0.00	0.86 ± 0.00	0.75 ± 0.00	0.88 ± 0.00
SCAR	0.76 ± 0.00	0.81 ± 0.00	0.80 ±0.00	0.83 ± 0.00	0.77 ± 0.00	0.80 ± 0.00	0.70 ± 0.00	0.81 ± 0.00	0.67 ± 0.03	0.77 ± 0.02
SpectACl	0.91 ± 0.03	0.90 ± 0.01	0.88 ± 0.04	0.89 ± 0.02	0.91 ± 0.00	0.95 ± 0.00	0.99 ± 0.00	1.00 ± 0.00	1.00 ±0.00	1.00 ±0.00
Ward Clust.	0.81	0.83	0.73	0.82	0.74	0.83	0.74	0.85	0.63	0.81
Birch	0.71	0.78	0.43	0.71	0.75	0.84	0.74	0.85	0.63	0.81
DEC	0.49 ± 0.03	0.63 ± 0.03	0.63 ± 0.06	0.76 ± 0.04	0.87 ± 0.03	0.92 ± 0.01	0.48 ± 0.11	0.68 ± 0.09	0.51 ± 0.03	0.73 ± 0.03
IDEC	0.69 ± 0.02	0.72 ± 0.02	0.63 ± 0.07	0.78 ± 0.03	0.86 ± 0.04	0.91 ± 0.01	0.75 ± 0.08	0.83 ± 0.04	0.75 ± 0.07	0.88 ± 0.03
DipDECK	0.74 ± 0.05	0.75 ± 0.02	0.72 ± 0.07	0.79 ± 0.04	0.73 ± 0.09	0.8 ± 0.03	0.81 ± 0.07	0.88 ± 0.03	0.75 ± 0.03	0.87 ± 0.01
DipEncoder	0.73 ± 0.05	0.76 ± 0.02	0.66 ± 0.08	0.77 ± 0.04	0.81 ± 0.06	0.86 ± 0.02	0.73 ± 0.10	0.85 ± 0.03	0.69 ± 0.06	0.83 ± 0.03
Mean Shift	0.88	0.83	0.89	0.83	0.91	0.84	0.95	0.87	0.95	0.88
DPCA	0.80	0.80	0.39	0.68	0.78	0.80	0.84	0.86	0.87	0.91
DCF	0.93	0.92	1.00	0.99	0.99	0.99	0.96	0.92	0.95	0.88
DDC	0.82	0.78	0.73	0.77	0.85	0.80	0.93	0.86	0.92	0.83
DBSCAN	0.97	0.95	0.90	0.90	0.90	0.94	0.99	1.00	1.00	1.00
OPTICS	0.98	0.97	0.94	0.97	0.85	0.93	1.00	1.00	1.00	1.00
HDBSCAN	0.95	0.94	0.98	0.97	0.92	0.92	0.99	1.00	1.00	1.00
MD-DBSCAN	0.97	0.96	1.00	1.00	1.00	1.00	0.99	0.99	1.00	1.00

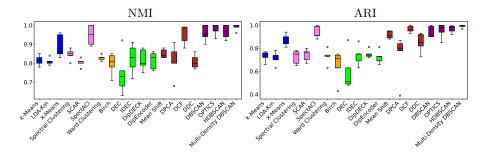


Fig. 5: NMI (left) and ARI (right) across all examined dimensionalities for the setting with "compact" intrinsic dimensionality. The coloration indicates the algorithm category as described in Appendix A.2.

Table 4: Best performing parameters (out of the evaluated ones) for different dimensionalities for the "compact"-Setting

dimensionality	2-dim	5-dim	10-dim	50-dim	100-dim
k-Means	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,
	n_init: 3	n_init: 2	n_init: 3	n_init: 7	n_init: 10
LDA-Km	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,
	n_init: 7	n_init: 8	n_init: 1	n_init: 1	n_init: 1
	n_clusters_init: 2,	n_clusters_init: 11,	n_clusters_init: 6,	n_clusters_init: 6,	n_clusters_init: 3,
X-Means	max_n_clusters: 14,	max_n_clusters: 12,	max_n_clusters: 10,	max_n_clusters: 10,	max_n_clusters: 10,
	n_split_trials: 3	n_split_trials: 10	n_split_trials: 10	n_split_trials: 2	n_split_trials: 3
	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,
Spectral Clust.	affinity: 'nearest_neighbors',	affinity: 'nearest_neighbors',	affinity: 'nearest_neighbors',	affinity: 'nearest_neighbors',	affinity: 'nearest_neighbors'
	nearest_neighbors: 19	nearest_neighbors: 30	nearest_neighbors: 190	nearest_neighbors: 100	nearest_neighbors: 200
	k: 10,	k: 10,	k: 10,	k: 10,	k: 10,
	nn: 75,	nn: 75,	nn: 30,	nn: 50,	nn: 100,
	alpha: 0.8,	alpha: 0.9,	alpha: 0.6,	alpha: 0.9,	alpha: 0.8,
SCAR	theta: 5000	theta: 1000	theta: 4000	theta: 4000	theta: 1000
	weighted: True,	weighted: True,	weighted: True,	weighted: True,	weighted: True,
	laplacian: 0,	laplacian: 0,	laplacian: 0,	laplacian: 0,	laplacian: 0,
	normalize: True	normalize: True	normalize: True	normalize: True	normalize: True
	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,	n_clusters: 10,
SpectACl	affinity: 'radius neighbors',	affinity: 'radius neighbors',	affinity: 'radius neighbors',	affinity: 'radius neighbors',	affinity: 'radius neighbors',
	epsilon: 0.08	epsilon: 0.25	epsilon: 0.12	epsilon: 0.15	epsilon: 0.15
W 100 :	n clusters: 10,	n clusters: 10,	n clusters: 10,	n clusters: 10,	n clusters: 10,
Ward Clust.	linkage: 'ward'	linkage: 'ward'	linkage: 'ward'	linkage: 'ward'	linkage: 'ward'
	n clusters: 10,	n clusters: 10,	n clusters: 10,	n clusters: 10,	n clusters: 10,
Birch	threshold: 0.01,	threshold: 0.01,	threshold: 0.01,	threshold: 0.01,	threshold: 0.01,
	branching factor: 2	branching factor: 2	branching factor: 2	branching factor: 2	branching factor: 2
	n clusters : 10,	n clusters : 10,			
DEC	alpha: 0.2,	alpha: 0.2,	alpha: 0.1,	alpha: 1.0,	alpha: 0.1,
-	embedding size: 2	embedding size: 2	embedding size: 10	embedding size: 5	embedding size: 5
	n clusters : 10,	n clusters : 10,	n clusters : 10,	n clusters: 10,	n clusters : 10,
	cluster loss weight: 0.01,	cluster loss weight: 1.0,	cluster loss weight: 0.1,	cluster loss weight: 1.0,	cluster loss weight: 1.0,
IDEC	alpha: 0.7,	alpha: 0.1,	alpha: 0.1,	alpha: 0.7,	alpha: 0.4,
	embedding size: 2	embedding size: 5	embedding size: 10	embedding size: 10	embedding size: 2
	cluster loss weight: 0.01,	cluster loss weight: 0.01,	cluster loss weight: 0.01,	cluster loss weight: 1.0,	cluster loss weight: 0.1,
	dip merge threshold: 0.85,	dip merge threshold: 0.95,	dip merge threshold: 0.75,	dip merge threshold: 0.95,	dip merge threshold: 0.95,
DipDECK	embedding size: 2,	embedding size: 5,	embedding size: 10,	embedding size: 2,	embedding size: 2,
Dipolicit	n clusters init: 15,	n clusters init: 15,			
	max n clusters: 20	max n clusters: 20			
	n clusters: 10,	n clusters: 10,	n clusters: 10,	n clusters: 10,	n clusters: 10,
DipEncoder	rec. loss weight: 10.0,	recloss_weight: 1000.0,			
Diplineodei	embedding size: 2	embedding size: 5	embedding size: 10	embedding size: 2	embedding size: 2
	cluster all: True,	cluster all: True,	cluster all: True,	cluster all: True,	cluster all: True,
Mean Shift	bandwidth: 0.06	bandwidth: 0.08	bandwidth: 0.11	bandwidth: 0.14	bandwidth: 0.14
	density threshold: 0.01,	density threshold: 0.01,	density threshold: 0.01,	density threshold: 0.01,	density threshold: 0.01,
DPCA	distance threshold: 0.07,	distance threshold: 0.09,	distance threshold: 0.1,	distance threshold: 0.15,	distance threshold: 0.15,
DICA	gauss cutoff: False	gauss cutoff: False	gauss cutoff: False	gauss cutoff: True	gauss cutoff: True
	k: 8,	k: 10,	k: 12,	k: 50,	k: 35,
DCF	beta: 0.97	beta: 0.95	beta: 0.97	beta: 0.99999	beta: 0.99999
	radius: 0.1,	radius: 0.1,	radius: 0.15,	radius: 0.18,	radius: 0.19,
DDC	merge: False	merge: True	merge: True	merge: True	merge: True
DBSCAN	epsilon: 0.015,	epsilon: 0.03,	epsilon: 0.05,	epsilon: 0.11,	epsilon: 0.11,
	min_samples: 3	min_samples: 2	min_samples: 8	min_samples: 2	min_samples: 2
	cluster_method: 'xi',	cluster_method: 'xi',	cluster_method: 'xi',	cluster_method: 'xi',	cluster_method: 'xi',
OPTICS	xi: 0.08,	xi: 0.03,	xi: 0.02,	xi: 0.08,	xi: 0.15,
	min_cluster_size: 150,	min_cluster_size: 100,	min_cluster_size: 100,	min_cluster_size: 10,	min_cluster_size: 10,
	predecessor_corr.: True	predecessor_corr.: True	predecessor_corr.: True	predecessor_corr.: True	predecessor_corr.: True
HDBSCAN	min_cluster_size: 12	min_cluster_size: 15	min_cluster_size: 3	min_cluster_size: 3	min_cluster_size: 2
	var: 3.0,	var: 1.8,	var: 1.5,	var: 1.2,	var: 1.2,
MD-DBSCAN	k: 10,	k: 10,	k: 10,	k: 25,	k: 30,
	min cluster size: 2	min cluster size: 2	min cluster size: 2	min cluster size: 5	min cluster size: 15

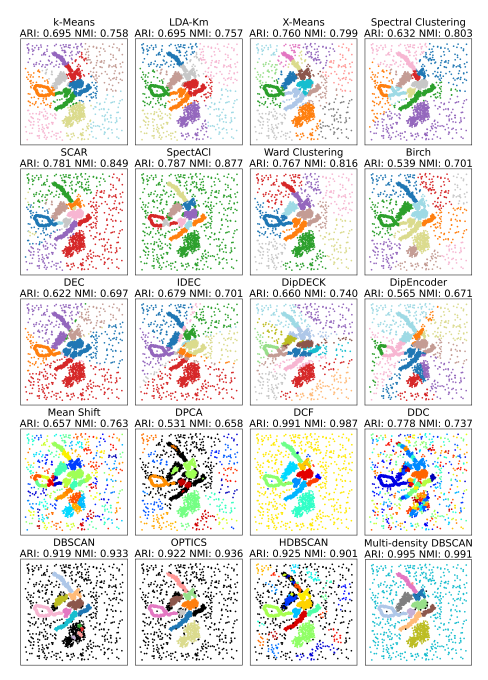


Fig. 6: State-of-the-art and fundamental clustering algorithms on the "mix"-setting dataset generated by DENSIRED with their corresponding performance

Table 5: Best performing parameters (out of the evaluated ones) for the "mix"-Setting

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set	11	$n \sigma$
JU	JUL.	112

dimensionality	2-dim
	n clusters: 10,
k-Means	n_init: 2
TD 4 17	n clusters: 10,
LDA-Km	n init: 1
	n_clusters_init: 7,
X-Means	max n clusters: 13,
11 Wicans	n split trials: 8
	n clusters: 10,
Spectral Clust.	affinity: 'nearest_neighbors',
Spectrar Clust.	nearest neighbors: 130
	k: 10,
	,
	nn: 15,
CCAD	alpha: 0.9,
SCAR	theta: 1000
	weighted: True,
	laplacian: 0,
	normalize: True
	n_clusters: 10,
SpectACl	affinity: 'radius_neighbors',
	epsilon: 0.05
Ward Clust.	n_clusters: 10,
ward Clust.	linkage: 'ward'
	n clusters: 10,
Birch	threshold: 0.025,
	branching factor: 200
	n clusters : 10,
DEC	alpha: 0.1,
DEC	embedding_size: 2
	n clusters: 10,
	cluster_loss_weight: 0.01,
IDEC	alpha: 0.3,
	embedding size: 2
	cluster_loss_weight: 0.01,
	dip_merge_threshold: 0.75,
Dis DECV	
DipDECK	embedding_size: 2,
	n_clusters_init: 15, max_n_clusters: 20
D: D 1	n_clusters: 10,
DipEncoder	reconstruction_loss_weight: 1000.0,
	embedding_size: 2
Mean Shift	cluster_all: True,
	bandwidth: 0.05
	density_threshold: 0.01,
DPCA	distance_threshold: 0.1,
	gauss_cutoff: True
DCF	k: 9,
DOF	beta: 0.9
DDC	radius: 0.1,
DDC	merge: False
DDCCAN	epsilon: 0.015,
DBSCAN	min_samples: 15
	cluster method: 'xi',
ODELGG	xi: 0.05,
OPTICS	min_cluster_size: 250,
	predecessor correction: True
HDBSCAN	min cluster size: 5
	var: 2.75,
MD-DBSCAN	k: 10,
D DDbonit	min cluster size: 5
	IIIII_CIGOUCI_DIZC. U

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