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**Highlights**

- Compare properties of different subspace clustering algorithms
- Introduce a new subspace clustering framework CSSub based on shared subspaces
- CSSub decouples the candidate subspace selection process from the clustering process
- CSSub enables different types of cluster definitions to be employed easily
- Provide different subspace generation and scoring methods

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# Grouping Points by Shared Subspaces for Effective Subspace Clustering

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## Abstract

Clusters may exist in different subspaces of a multidimensional dataset. Traditional full-space clustering algorithms have difficulty in identifying these clusters. Various subspace clustering algorithms have used different subspace search strategies. They require clustering to assess whether cluster(s) exist in a subspace. In addition, all of them perform clustering by measuring similarity between points in the given feature space. As a result, the subspace selection and clustering processes are tightly coupled. In this paper, we propose a new subspace clustering framework named *CSSub* (Clustering by Shared Subspaces). It enables neighbouring core points to be clustered based on the number of subspaces they share. It explicitly splits candidate subspace selection and clustering into two separate processes, enabling different types of cluster definitions to be employed easily. Through extensive experiments on synthetic and real-world datasets, we demonstrate that *CSSub* discovers non-redundant subspace clusters with arbitrary shapes in noisy data; and it significantly outperforms existing state-of-the-art subspace clustering algorithms.

*Keywords:* Subspace clustering, Shared subspaces, Density-based clustering.

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## 1. Introduction

Clustering detects groups of similar points while separating dissimilar points into different groups [1]. Many clustering algorithms have been studied in the research community over the past few decades. Clustering is a fundamental data analysis technique for data mining and knowledge discovery and has been applied to various fields such as engineering, medical and biological sciences, social sciences and economics.

Traditional “full-space clustering” algorithms become ineffective when clusters exist in different subspaces<sup>1</sup> as many irrelevant attributes may cause similarity measurements used by the algorithms

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<sup>1</sup>By subspace we intend a linear subspace over a subset of the original dimensions. Moreover, we say that a “cluster exists in a subspace” if the cluster is defined as a region within the subspace, and thus dimensions not included in the

to become unreliable [1]. To tackle this problem, the similarity of points should be assessed within  
 10 subspaces (over a subset of relevant attributes).

Subspace clustering aims to discover clusters which exist in different subspaces [2]. It works by  
 combining subspace search and clustering procedures. The number of possible axis-parallel subspaces  
 is exponential in the dimensionality and the number of possible arbitrarily oriented (non axis-parallel)  
 subspaces is infinite. To deal with the huge number of possible subspaces, subspace clustering algo-  
 15 rithms usually rely on heuristics for subspace search, e.g., top-down search [3] and bottom-up search  
 [4]. All these existing subspace clustering algorithms perform clustering by measuring the similarity  
 between points in the given feature space, and the subspace selection and clustering processes are  
 tightly coupled.

In this paper, we focus on clustering in axis-parallel subspaces. We contribute a new subspace  
 20 clustering framework, named *CSSub* (Clustering by Shared Subspaces)<sup>2</sup>, which has the following  
 three unique features:

1. *CSSub* groups points by their shared subspaces. It performs clustering by measuring the similar-  
 ity between points based on the number of subspaces they share. This enables *CSSub* to detect  
 non-redundant/non-overlapping subspace clusters directly by running a clustering method only  
 25 once. In contrast, many existing subspace clustering algorithms need to run a chosen cluster-  
 ing method for each subspace; and this must be repeated for an exponentially large number of  
 subspaces. As a consequence, it produces many redundant subspace clusters.
2. *CSSub* decouples the candidate subspace selection process from the clustering process. By  
 explicitly splitting them into independent processes, it enables candidate subspaces to be selected  
 30 independent of the clustering process—eliminating the need to repeat the clustering step a large  
 number of times. In contrast, many existing subspace clustering algorithms which have tightly-  
 coupled processes must rely on an anti-monotonicity property to prune the search space.
3. The decoupling approach has an added advantage that allows different types of cluster definitions  
 to be employed easily. The time cost of the entire process is dominated by the subspace scoring  
 35 function. We show that by changing the cluster definition such that subspaces can be evaluated  
 with a linear scoring function, enables the runtime of *CSSub* to be reduced from quadratic time  
 to linear time. A similar change is difficult, if not impossible, for existing algorithms because of  
 the tightly coupled processes.

We present an extensive empirical evaluation on synthetic and real-world datasets to demonstrate

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subspace are irrelevant and can be considered noise with respect to the cluster.

<sup>2</sup>The source code of *CSSub* can be obtained at <https://sourceforge.net/projects/subspace-clustering/>.

40 the effectiveness of *CSSub*. The experiments show that *CSSub* discovers subspace clusters with arbitrary shapes in noisy data; and it significantly outperforms existing state-of-the-art subspace clustering algorithms. In addition, *CSSub* has only one parameter  $k$  (the number of clusters) which needs to be manually tuned, other parameters can be automatically set based on a heuristic method.

The rest of this paper is organised as follows. We provide an overview of subspace clustering algorithms and related work in Section 2. Section 3 discusses key weaknesses of existing bottom-up  
45 subspace clustering algorithms. Section 4 details the subspace clustering framework based on the shared subspaces and presents a density-based approach for subspace scoring. Section 5 presents the algorithms for *CSSub*. In Section 6, we empirically evaluate the performance of the proposed algorithms on different datasets. Discussion and the conclusions are provided in the last two sections.

## 50 2. Related Work

The key task of clustering in subspaces is to develop appropriate subspace search heuristics [2]. There are two basic techniques for subspace search, namely top-down search [3] and bottom-up search [4]. Different subspace clustering algorithms have been proposed based on these two search directions [5, 6, 7, 8, 9]. Search strategies can be further subdivided into systematic and non-systematic search  
55 techniques as discussed below.

### 2.1. Systematic Subspace Search

Systematic search strategies for subspace clustering can be top down [3] or bottom up [4]. Searching all possible subspaces generates  $2^d - 2$  candidates (where  $d$  is the number of distinct attributes). To avoid having to perform an exhaustive search, the two systematic search strategies utilise certain  
60 criteria to prune the search space.

#### 2.1.1. Bottom-up Methods

The bottom-up subspace search strategy uses the *Apriori* algorithm [5]. It starts from all one-dimensional subspaces and then searches for higher dimensional subspaces progressively. This strategy relies on the anti-monotonicity property to prune the search space: if a candidate subspace in a lower  
65 dimensional space has no clusters or is filtered out according to some other criteria, then its projection onto a higher dimensional space will not be traversed.<sup>3</sup>

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<sup>3</sup>To be precise, the anti-monotonicity property is a property of density-based clustering approaches only, where points can be considered either “high density and therefore within a cluster” or “low density and therefore noise”. Since density monotonically reduces with increased dimension, if a fixed density threshold is used to define points as high density then the property will be anti-monotone (in the set of attributes), i.e., a low-density point in a subspace with a set of attributes is still a low-density point in a higher dimensional subspace which contains these attributes.

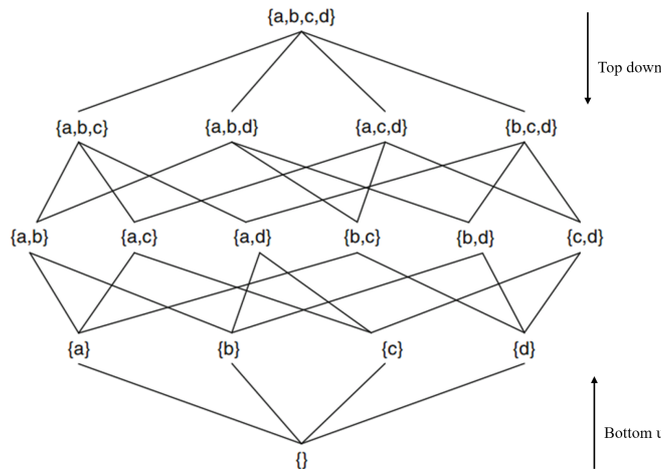


Figure 1: The lattice of candidate subspaces from attributes  $\{a, b, c, d\}$ .

CLIQUE [4] uses a grid-based structure to investigate dense units at  $k - 1$  dimensional subspace to produce the possible set of  $k$ -dimensional units that might contain dense units. MAFIA [10] is an extension of CLIQUE that uses a variable-width grid-based density estimator in each subspace in order to improve efficiency and quality. ENCLUS [11] is similar to CLIQUE, but it uses entropy rather than density to measure the “clusterability” of a subspace, i.e., subspaces having lower entropy normally contain more meaningful clusters. ENCLUS uses the same bottom-up fashion of CLIQUE for finding subspace clusters. However, the grid-based algorithm cannot be applied on high-dimensional datasets as the number of cells in the grid grows exponentially with the dimensionality of data.

SSCS [12] is based on the monotonicity property of density-based subspace clustering to tackle the exponential subspace search problem. Instead of performing step-by-step scheme of the *Apriori* algorithm, it proposes a best-first style such that it directly steers to high dimensional subspaces from low dimensional subspaces with successive information gathering about dense subspace regions identified by hyper-rectangles. It only outputs the most interesting clusters in a few high-dimensional subspaces for non-redundant clustering.

P3C [13] is a grid-based algorithm relying on statistical significance tests to detect subspace clusters. P3C defines significant regions as regions containing significantly more points than expected under a uniform-distribution assumption over a particular subspace. P3C first detects significant regions in individual dimensions by starting with intervals and merging adjacent significant regions together to form cluster cores. It then searches for higher dimensional regions using a bottom-up search strategy. Finally, P3C produces a matrix that records the probability of each point belonging to each cluster. P3C reports the results of the final clustering using an EM-like clustering procedure [14]. The advantage is that P3C can be used for both numerical and categorical data. Like other density-based algorithms,

P3C can detect arbitrarily shaped clusters and is robust to noise as well. The main disadvantage of  
 90 P3C is that its performance depends on the detected one-dimensional clusters. If clusters have low  
 density in their one-dimensional projections, there will not be enough statistical evidence for further  
 aggregation.

Although the grid-based algorithms provide efficient clustering models, they cannot be applied to  
 high-dimensional datasets, as the number of cells in the grid grows exponentially with the dimen-  
 95 sionality of data. The cluster shape detected by grid-based clustering is a polygon (which can be an  
 arbitrary shape for a high grid granularity) in the corresponding subspace where the space is divided  
 in cells by a grid. Their accuracy and efficiency depend on the granularity and the positioning of the  
 grid [2].

In addition, SUBCLU [15] and FIRES [16] are similar to CLIQUE, but apply different methods  
 100 for clustering points. SUBCLU applies DBSCAN rather than grid-based clustering on each subspace  
 candidate and outputs all detected clusters when searching for higher dimensional subspaces progres-  
 sively. FIRES is a general framework for efficient subspace clustering. It runs a clustering algorithm  
 on individual attributes separately in order to find “base clusters” and then utilises an approximation  
 method for merging base clusters iteratively in order to find maximal attribute combinations in sub-  
 105 spaces with higher dimensionality. After that, some methods can be used to refine the final clustering  
 results. Its performance highly depends on the pruning and refinement steps [16].

Generally, clustering algorithms based on bottom-up searches normally result in overlapping/redundant  
 clusters, such that a point can belong to many clusters simultaneously, making the clusters difficult to  
 interpret. SUBCLU and the grid-based algorithms use a global density threshold, which leads to a bias  
 110 against higher dimensional subspaces, i.e., a tighter threshold can filter noise well in low-dimensional  
 subspaces but will lose clusters in higher dimensional subspaces. P3C can avoid this bias by using a  
 threshold based on a statistical significance test, which can be treated as an adaptive density threshold.  
 However, these bottom-up algorithms will take an extremely long time to discover subspace clusters  
 if those clusters are high-dimensional, i.e., if the subspace they occupy spans many or most of the  
 115 original attributes.

### 2.1.2. Top-down Methods

In order to find the best non-overlapping partitioning of the data into subspaces, most top-down  
 methods aim to group points such that each point belongs to only one cluster and each cluster is  
 assigned to a specific subspace [3]. This search method determines the subspace of a cluster starting  
 120 from the full-dimensional space and then removes the irrelevant attributes iteratively. These methods  
 usually evaluate each attribute for each point or cluster based on the “locality assumption” [6], i.e.,  
 the subspace of each cluster can be learnt from the local neighbourhood of cluster members. Note that

in some papers, subspace clustering with a top-down method is also called “projected clustering” [2].

In order to find non-overlapping data partitionings in subspaces, most top-down based subspace  
 125 clustering algorithms aim to group points such that each point belongs to one cluster only; and each  
 cluster is assigned to a specific subspace [3]. A top-down algorithm begins the search from the full-  
 dimensional space and then removes the irrelevant attributes iteratively to find the target subspace.

PROCLUS [3] is a  $k$ -medoids type [17] clustering algorithm. It first randomly selects  $k$  potential  
 cluster centres from the data sample and then iteratively refines them. In each loop of the iterative  
 130 process, for each medoid, a set of dimensions is chosen by minimising the standard deviation of the  
 distances of all points in the neighbourhood of the medoids to the corresponding medoid along every  
 dimension. Points are then regrouped to their closest medoid based on the identified subspace of each  
 medoid. In the final step of each loop, medoids are updated based on the points currently assigned  
 to them. The iterative process continues until the clustering quality cannot be improved. The points  
 135 which are too far away from their closest medoids are identified as noise. The output is  $k$  groups of  
 non-overlapping points and a set of noise points. Each cluster has a globular shape in its identified  
 subspace. The performance of PROCLUS is not stable since the initial medoids are randomly selected  
 and are sensitive to irrelevant attributes in the full-dimensional space.

PreDeCon [18] is a top-down clustering algorithm which relies on DBSCAN [19]. For each point,  
 140 this algorithm calculates a specialised subspace distance based on the concept of subspace preference,  
 using it for density estimation and linking points. An attribute is relevant to the “subspace preference”  
 of a point if the variance of data in the point’s  $\epsilon$ -neighbourhood is considered smaller than a threshold.  
 It then links neighbouring points based on their density distribution on the weighted attributes. The  
 downside of this approach is that there are three parameters that need to be manually set for density  
 145 estimation [18]. Since it does not identify the subspaces where clusters exist but only weights attributes  
 for each cluster, it is referred to as a “soft” subspace clustering algorithm (while others are called “hard”  
 subspace clustering algorithms).

DOC [20] mixes the bottom-up approach for cluster detection and the top-down approach for  
 iterative improvement in order to reduce redundancy. It uses a fixed density threshold with fixed  
 150 side-length hypercubes to identify clusters.

## 2.2. Non-systematic Subspace Search

There are some subspace clustering algorithms which do not rely on a systematic search strategy  
 but pursue heuristic approaches. For example, STATPC [21] formulates the search for statistically  
 significant regions as subspace clusters with a grid-based structure. It defines the search as an optimi-  
 155 sation problem and proposes a heuristic solution to avoid an exhaustive search.

In addition, there is rich literature on soft subspace clustering derived from  $k$ -means type [22] clustering. Different from selecting subspaces for each cluster, this kind of algorithms learn weight vectors over attributes with different objective functions and optimisation methods, and then incorporate these vectors for distance calculation for the  $k$ -means algorithm.

160 Generally, soft subspace clustering algorithms can be classified into three categories [23], i.e., conventional (all clusters share the same subspace, e.g., C- $k$ -means [24]), independent (each cluster has an independent subspace, e.g., LAC [25], EWKM [26] and FSC [27]) and extended (enhanced performance for specific purposes, e.g., ESSC [28] and FG- $k$ -means [29]). They also can be divided into soft subspace fuzzy clustering (e.g., FSC [27]) and soft subspace hard clustering (e.g., the five algorithms, 165 apart from FSC, mentioned before), depending on whether each data object belongs to only one cluster or multiple clusters with different degrees [30, 31].

LAC, for example, starts with  $k$  centroids and  $k$  sets of  $d$  random weights, and approximates a set of  $k$  Gaussians by adapting the weights [25]. Soft subspace clustering algorithms can be seen as performing some type of normalisation of attributes for each cluster. In addition,  $k$ -means-based soft 170 subspace clustering usually requires one parameter  $k$  as the number of clusters while other parameters can be set to default values [29].

EWKM is an entropy weighting subspace clustering algorithm which simultaneously maximises the negative weight entropy and minimises the within-cluster dispersion [26]. It gives different weights to attributes by utilising the weight entropy in the objective function. Therefore, clusters may have 175 different selected attributes based on their weights.

Both ESSC and FG- $k$ -means are based on EWKM method for specific purposes. ESSC integrates both within-cluster compactness and between-cluster separation information [28]. FG- $k$ -means introduces multi-feature weighting for co-learning and multi-view learning [29]. They have better performance than EWKM, but FG- $k$ -means and ESSC introduce 1 and 2 more parameters, respectively.

180 It is worth mentioning that there are many algorithms related with bi-clustering, co-clustering and pattern-based clustering [1]. They find clusters based on the pattern in the data matrix, and to cluster both objects and attributes simultaneously. They are usually aimed for genes and microarray datasets, and can only detect special types of subspace clusters [2]. They are not the focus of this paper.

### 2.3. Summary

185 Figure 2 compares properties of different subspace clustering algorithms. All existing subspace clustering tightly coupled their subspace selection process and clustering process; clustering definition cannot be changed easily. Note that the weights search in soft subspace clustering is also tightly coupled with the clustering process based on some optimisation functions.  $k$ -medoids and  $k$ -means type clustering algorithms usually cannot detect non-globular shaped clusters [2]. Soft subspace clustering,

Figure 2: Properties of subspace clustering algorithms and the proposed *CSSub* framework.

Algorithm	Subspace search strategy	Cluster subspace assignment	Cluster type	Arbitrarily shaped clusters	Non-overlapping clusters	Adaptive density clusters	Handling noise	Easy parameter setting	Clustering without original attributes
CLIQUE	Bottom-up	Hard	Grid	✓			✓		
MAFIA	Bottom-up	Hard	Grid	✓			✓		
ENCLUS	Bottom-up	Hard	Grid	✓			✓		
SSCS	Bottom-up	Hard	Density	✓	✓		✓		
P3C	Bottom-up	Hard	Grid	✓	✓	✓	✓	✓	
FIRES	Bottom-up	Hard	Flexible	✓		✓	✓		
SUBCLU	Bottom-up	Hard	Density	✓			✓		
PROCLUS	Top-down	Hard	$k$ -medoids		✓		✓	✓	
PreDeCon	Top-down	Soft	Density	✓	✓		✓		
DOC	Hybrid	Hard	Grid	✓	✓				
STATPC	Heuristic	Hard	Grid	✓		✓	✓		
C- $k$ -means	Optimisation	Soft	$k$ -means		✓			✓	
FSC	Optimisation	Soft	$k$ -means		✓			✓	
LAC	Optimisation	Soft	$k$ -means		✓			✓	
EWKM	Optimisation	Soft	$k$ -means		✓			✓	
ESSC	Optimisation	Soft	$k$ -means		✓			✓	
FG- $k$ -means	Optimisation	Soft	$k$ -means		✓			✓	
<i>CSSub</i>	Non-search	Hard	Flexible	✓	✓	✓	✓	✓	✓

190 especially those based on  $k$ -means, cannot handle noise. Most density-based algorithms have more than 2 critical parameters for defining clusters, which are very difficult to set in practice.

This paper focuses on hard subspace and non-overlapping clustering. It is important to mention that the *CSSub* framework does not rely on any subspace search strategies or optimisation. Since *CSSub* clusters points based on the number of subspaces they shared, original attributes are not used  
 195 for the similarity calculation in the clustering stage. In contrast, all the other algorithms rely on the original attributes.

### 3. Key weaknesses of existing bottom-up subspace clustering algorithms

The majority of density-based subspace clustering algorithms rely on a bottom-up search strategy and use anti-monotonicity of the density as a means to reduce the search space. This approach has  
 200 two weaknesses:

1. The approach tightly couples the candidate subspace selection process with the clustering process,

i.e., it must run a clustering method for each subspace; and there are an exponentially large number of subspaces. As a result, many redundant subspace clusters are produced during the search process. This is a typical problem of using bottom-up search to find subspaces; and it has its origin in finding frequent item-sets [2]. While there are techniques to reduce the redundancy after the candidate subspaces have been generated, the approach is inefficient and ineffective in producing non-redundant subspace clusters.

2. The requirement to use a single (fixed) density threshold in order to make use of the anti-monotonicity property to prune the search space. Because density is a dimensionality biased measure [32], any global threshold selected will bias the search for subspace clusters toward low dimensional subspaces. While it is possible to use an adaptive density threshold or adaptive neighbourhood radius to detect more core points in higher dimensional subspaces [33], these methods cannot be directly deployed on density-based clustering algorithms. Incorporating these adaptive methods often involve ad hoc adjustments and with additional parameters.

To overcome the above weaknesses, we propose to use a different approach which does not rely on search and needs to run the clustering method only once to produce non-redundant subspaces.

#### 4. Clustering by Shared Subspaces

In this section, we present an effective subspace clustering framework in order to overcome the weaknesses of existing bottom-up subspace clustering algorithms.

##### 4.1. Definitions

Let  $D$  be a dataset of  $n$  points of  $d$  attributes, represented in a  $n \times d$  matrix. A subspace cluster  $C_i$  is a submatrix of  $D$  with  $|s_i| < d$  attributes, and  $|C_i| \leq n$ . We denote  $S = \{s_1, s_2, \dots, s_m\}$  as the set of subspaces to be explored. Let  $\pi_s(x)$  denote the point  $x \in D$  projected on a subspace  $s \in S$ . Let the set of  $k$  subspace clusters in  $D$  be  $C = \{C_1, C_2, \dots, C_k\}$ .

**Definition 1.** Point  $x$  is an  $\alpha$ -Core point in subspace  $s$  if  $x$  has a high score value based on a scoring function  $\alpha(\cdot)$  in  $s$ , i.e.,  $(\alpha_s(x) > \tau_s) \leftrightarrow \alpha\text{-Core}_s(x)$ . Subspace  $s$  is thus a candidate subspace (core-point subspace) for  $x$  in which a cluster containing  $x$  may exist.

Note that  $\tau_s$  depends on subspace  $s$ ; and it is not a global threshold for all subspaces. Noise points are non- $\alpha$ -Core points in all subspaces considered.

**Definition 2.** Similarity by *shared subspaces*: the similarity between  $x_i$  and  $x_j$  is defined as the number of shared subspaces in which both  $x_i$  and  $x_j$  are  $\alpha$ -Core.

For example, if density is used as the score  $\alpha(\cdot)$ , then the similarity between  $x_i$  and  $x_j$  is defined as the number of subspaces in which they are both in dense locations.

Let  $A(x) = \{s \in S \mid \alpha\text{-Core}_s(x)\}$  be the set of subspaces where point  $x$  is an  $\alpha$ -Core point. As each point is  $\alpha$ -Core in a set of subspaces which is different from that of another point, the similarity is formulated through normalisation as follows:

$$\text{sim}(x_i, x_j) = \frac{|A(x_i) \cap A(x_j)|}{|A(x_i) \cup A(x_j)|}$$

Note that the above formulation is the Jaccard similarity [34]; it can be interpreted as the higher the probability that two  $\alpha$ -Core points are in shared subspaces, the more similar the two points are.

**Definition 3.** Cluster  $C$  exists in a set of shared subspaces  $S$  if  $\forall_{i \neq j, x_i, x_j \in C} \text{sim}(x_i, x_j) \geq \beta_C$ , where  $\beta_C$  is a threshold for cluster  $C$ .

Note that  $\beta_C$  varies for each cluster and it can be determined automatically if a certain clustering algorithm is employed, e.g., a partitioning-based algorithm  $k$ -medoids. Hereafter we denote  $\alpha$ -Core points as “core points”.

#### 4.2. Clustering Using Shared Subspaces

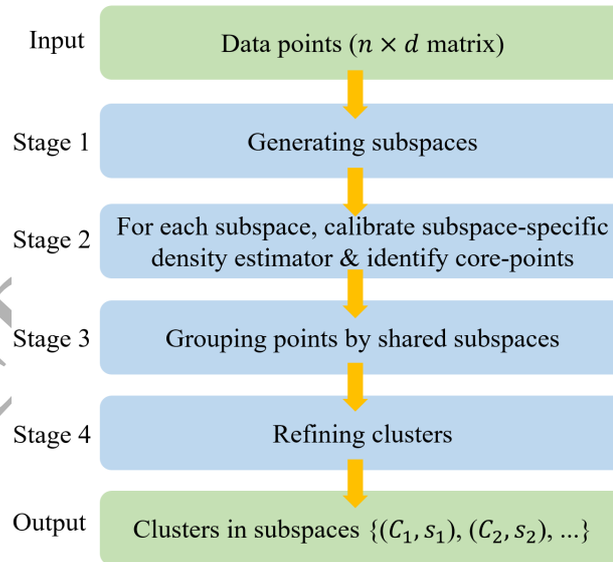


Figure 3: A conceptual overview of the *CSSub* framework for subspace clustering.

A conceptual overview of the *CSSub* framework is shown in Figure 3. There are four stages. In Stage 1, we first generate an initial set of subspaces for assessing. Then each point is examined whether it is a core point in each of the subspaces in Stage 2. Candidate subspaces are selected if they have

245 core points. Non-core points in all subspaces are identified as noise. In Stage 3, by using a similarity measure based on shared subspaces, such as the Jaccard similarity, an existing clustering algorithm can be employed to perform clustering on the points defined using the candidate subspaces identified in Stage 2. In Stage 4, some methods are applied to refine these clusters, e.g., finding the best subspace for each cluster to produce non-overlapping clusters.

#### 250 4.2.1. Scoring Function

*CSSub* can produce different types of clusters depending on the scoring function employed. We first use a density scoring function to demonstrate the ability of *CSSub* to detect arbitrarily shaped clusters. We then show that a different score can be easily used in the framework in Section 6.4.

To use a density function to identify core points, a density estimator is required. We employ the  $\epsilon$ -neighbourhood density estimator [19] as follows:

$$\widehat{pdf}_s^\epsilon(x) = \frac{1}{nV_d^\epsilon} \sum_j \mathbf{1}[\|\pi_s(x) - \pi_s(x_j)\| \leq \epsilon] \quad (1)$$

where  $\mathbf{1}[\cdot]$  denotes the indicator function and  $\pi_s(x)$  is point  $x \in D$  projected on subspace  $s \in S$ , and 255  $V_d^\epsilon \propto \epsilon^d$  is the volume of a  $d$ -dimensional ball of radius  $\epsilon$ . As the volume is constant for all points, it is usually omitted in actual computations.

Since data become sparse as the dimensionality of subspace increases, a global  $\epsilon$  is not appropriate, i.e., using a relatively very small or very large  $\epsilon$  may get a uniform density distribution for each point in a subspace. In order to maximally differentiate points in terms of their estimated density values in a 260 subspace  $s$ , we use an adaptive  $\epsilon_s$  setting to obtain the maximum variance of the normalised densities such that:  $\epsilon_s = \arg \max_{\epsilon \in R} \sum_j (p_s^\epsilon(x_j) - \frac{1}{n} \sum_k p_s^\epsilon(x_k))^2$ , where  $p_s^\epsilon(x) = \widehat{pdf}_s^\epsilon(x) / \sum_y (\widehat{pdf}_s^\epsilon(y))$  is the normalised density of  $x$  in subspace  $s$  given  $\epsilon$ . In our implementation, a fix number  $g$  of possible  $\epsilon$  values is examined to obtain the best value  $\epsilon_s$  for each subspace.

Let  $p_s^{\epsilon_s}(x)$  be the density score of point  $x$  in subspace  $s \in S$ . We use  $p_s^{\epsilon_s}(\cdot)$  as the subspace scoring 265 function in Stage 2. Subspace  $s$  is a selected candidate subspace for  $x$  if  $(p_s^{\epsilon_s}(x) > \tau_s)$ . Note that  $p_s^{\epsilon_s}(\cdot)$  is used in place of  $\alpha_s(\cdot)$  in Definition 1 to determine core points in subspace  $s$ . Here we propose a heuristic method that set  $\tau_s$  to the average score in each subspace as the default parameter, i.e.,  $\tau_s = \frac{1}{n} \sum_{x \in D} \alpha_s(x) = \frac{1}{n} \sum_{x \in D} p_s^{\epsilon_s}(x)$ . This means that every core point has a density larger than the average density in the subspace. Thus, parameters  $\epsilon_s$  and  $\tau_s$  in this density-based scoring function are 270 set automatically.

#### 4.2.2. Grouping Algorithm

A traditional clustering algorithm can be used to group points based on their Jaccard similarities. We focus on discovering non-overlapping clusters  $C$  such that  $\forall_{i \neq j, C_i, C_j \in C} C_i \cap C_j = \emptyset$ . In this paper, we use  $k$ -medoids algorithm [17] for finding  $k$  optimal non-overlapping partitions.

**Algorithm 1** *Clustering\_by\_Shared\_Subspaces*( $D, k$ )**Input:**  $D$ : input data ( $n \times d$  matrix);  $k$ : number of clusters.**Output:**  $\{C_i, s_i\}_{i=1}^k$ : subspace clusters; *noise*: noise points.

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```

/* Stage 1: */
1:  $d_{max} \leftarrow \max \dim \text{ s.t. } \sum_{i=1}^{\dim} \binom{d}{i} < n$ ;
2:  $S \leftarrow \text{generate\_subspaces}(D, d_{max})$ ;
/* Stage 2: */
3: for  $s \in S$  do
4:    $\epsilon_s \leftarrow \arg \max_{\epsilon \in R} \sum_j (p_s^\epsilon(x_j) - \frac{1}{n} \sum_k p_s^\epsilon(x_k))^2$ ;
5:    $\text{core}_{s,j} \leftarrow \mathbf{1}[p_s^{\epsilon_s}(x_j) > \frac{1}{n} \sum_k p_s^{\epsilon_s}(x_k)] \quad \forall_{j=1}^n$ ;
6: end for
7:  $\text{noise} \leftarrow \{x_j \in D \mid \forall_{s \in S} \text{core}_{s,j} = 0\}$ ;
8:  $\text{core} \leftarrow \forall_{s,j} \text{remove } \text{cores}_{s,j} \text{ if } x_j \in \text{noise}$ ;
/* Stage 3: */
9:  $\{C_i\}_{i=1}^k \leftarrow k\_medoids(\text{core}, k)$ ;
/* Stage 4: */
10: for  $i \in \{1, \dots, k\}$  do
11:    $\text{shared}_i = \{s \in S \mid \exists_{x_j \in C_i} \text{core}_{s,j} = 1\}$ ;
12:    $s_i \leftarrow \arg \max_{s \in \text{shared}_i} \sum_{x_j \in C_i} \text{core}_{s,j}$ ;
13: end for
14: return  $\{C_i, s_i\}_{i=1}^k$ ; noise;

```

---

275  $k$ -medoids is more resilient to noise and outliers than  $k$ -means. Instead of using the mean value to represent a cluster,  $k$ -medoids [17] chooses one data points  $c_i \in C_i$  as the centre (medoid) for a cluster. A medoid is a point with the minimal average distance to all the points in the cluster. Thus, the best medoid for each cluster is found by minimising the sum of the dissimilarities between each point in the cluster and a candidate medoid.

## 280 5. Algorithms in *CSSub*

In this section, we provide the algorithms for the proposed *CSSub* framework shown in Figure 3. The stages shown in Algorithm 1 correspond to the stages in Figure 3.

285 The first stage in Algorithm 1 generates the set of subspaces by enumeration to the largest subspace dimensionality  $d_{max} \geq 1$  such that  $|S| = \sum_{i=1}^{d_{max}} \binom{d}{i} = m < n$ . Here we set the number of candidate subspaces less than the data size is to keep the quadratic time complexity. Different ways to generate the initial set of subspaces may be used instead in the framework, discussed in Section 7.1.

**Algorithm 2**  $k\_medoids(core, k)$ **Input:**  $core$ : core-point subspace matrix;  $k$ : number of desired clusters.**Output:**  $C$ : a set of clusters.

- 1: Randomly select  $k$  data points from  $core$  as the medoids;
- 2: Each data point from  $core$  is assigned to the closest medoid to form clusters  $C_i, i = 1, \dots, k$  based on the Jaccard similarity;
- 3: Revise the medoid in each cluster which yields the maximum sum of the similarity between each point in the cluster and the medoid, i.e., for each cluster  $i$ , find medoid  $M_i = \arg \max_{y \in C_i} \sum_{x \in C_i} sim(x, y)$ ;
- 4: Repeat steps 2 and 3 until there is no change in point reassignments;
- 5: **return**  $C = \{C_1, \dots, C_k\}$ ;

In Stage 2 of Algorithm 1, the output  $core$  is a binary matrix, which indicates whether a point is a core point in a subspace. It also outputs the noise points which are not core points in any subspaces assessed; they are excluded for clustering.

290 In Stage 3 of Algorithm 1, we use a  $k$ -medoids algorithm based on the PAM (Partitioning Around Medoids) algorithm [17] to cluster the data on the matrix  $core$  using the Jaccard similarity. The  $k$ -medoids algorithm is shown in Algorithm 2.

For each detected cluster  $C_i \subset C$  at the end of Stage 3 in Algorithm 1, there exists a set of shared subspaces  $shared_i = \{s \in S \mid \alpha-Core_s(x) \wedge (x \in C_i)\}$ .

295 In Stage 4 of Algorithm 1, we refine the clustering result of  $k$ -medoids in order to find the best subspace for each cluster. The final subspace for  $C_i$  is the one in  $shared_i$  which covers the largest number of points in  $C_i$ , i.e.,  $s_i = \arg \max_{s \in shared_i} |\{x \in C_i \mid \alpha-Core_s(x)\}|$ .

The final clustering outputs are the  $k$  clusters:  $\{(C_1, s_1), (C_2, s_2), \dots, (C_k, s_k)\}$ ; and noise points identified in Stage 2.

300 It is interesting to note that in using  $k$ -medoids to group points by shared subspaces, the algorithm implicitly sets  $\beta_{C_i}$  in Definition 3 automatically for each cluster  $C_i$  having medoid  $M_i$  as follows:

$$\beta_{C_i} = \min_{x \in C_i} sim(x, M_i).$$

$CSSub$  requires only one parameter  $k$  to be set manually, and has the total time complexity of  $O(gn^2 d^{d_{max}})$ .

## 305 6. Empirical Evaluation

This section presents experiments designed to evaluate the performance of  $CSSub$ . We selected seven state-of-the-art non-overlapping subspace clustering algorithms for comparison: a top-down subspace clustering algorithm PROCLUS [3], a bottom-up subspace clustering algorithm P3C [13] and

five soft subspace clustering algorithms LAC [25], EWKM [26], FSC [27] ESSC [28] and FG- $k$ -means  
 310 [29]. In addition, we include a full-space clustering algorithms  $k$ -medoids [17] in order to judge the  
 effectiveness of subspace clustering for real world tasks. Because it is difficult to assess the clustering  
 performance for algorithms which produce many redundant clusters, this type of subspace clustering  
 algorithms is omitted in the comparison.

We used 3 synthetic datasets (2T, S1500 and D50) and 12 real-world datasets<sup>4</sup> to ascertain the  
 315 ability of different subspace clustering algorithms in handling different kinds of datasets. All datasets  
 are normalised using the *min-max* normalisation. The properties of these datasets are presented in  
 Table 1.

Table 1: The properties of 15 datasets. The first 3 datasets are synthetic datasets and the rest are real-world datasets.

Datasets	#Points	#Dimensions	#Clusters
2T	1600	4	2
S1500	1595	20	12
D50	1596	50	13
Glass	214	9	6
Liver	345	6	2
ILPD	579	9	2
Musk	476	166	2
Sonar	208	60	2
Ionosphere	351	33	2
Spam	4601	57	2
Iris	150	4	3
Wine	178	13	3
Dermatology	358	34	6
WDBC	596	30	2
Breast	699	9	2

The 2T synthetic dataset contains two T-shaped clusters embedded in a four-dimensional space,  
 while the S1500 and D50 synthetic datasets have 12 and 13 Gaussian clusters, respectively, embedded  
 320 in different subspaces. Every irrelevant attribute in these synthetic datasets is uniformly distributed  
 between 0 and 1. Figure 4 and Figure 5 show the data distributions of the 2T and S1500 datasets in  
 two different subspaces, respectively. The D50 dataset has 50 attributes and more irrelevant attributes  
 than the S1500 dataset.

PROCLUS, FG- $k$ -means, LAC and P3C were implemented in Weka, and all other algorithms were  
 325 implemented in Matlab<sup>5</sup>. The experiments were run on a machine with eight cores (Intel Core i7-  
 7820X 3.60GHz) and 32GB memory. All datasets were normalised using the *min-max* normalisation

<sup>4</sup>Synthetic datasets S1500 and D50 are from [8]. All real-world datasets are from UCI Machine Learning Repository [35].

<sup>5</sup>The source codes of PROCLUS and P3C are from the paper [8]. FG- $k$ -means and LAC source codes are from the paper [29]. Other soft subspace clustering algorithms were from the paper [23].

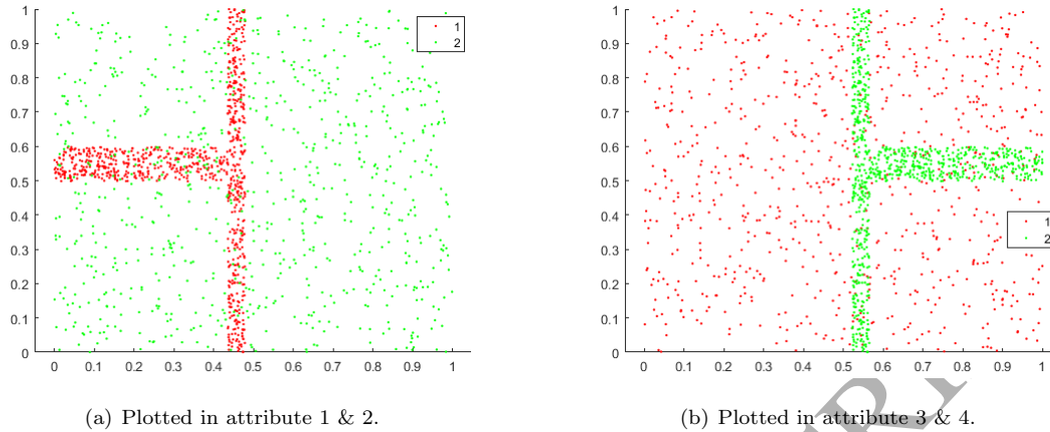


Figure 4: Distributions of the 2T dataset in two subspaces.

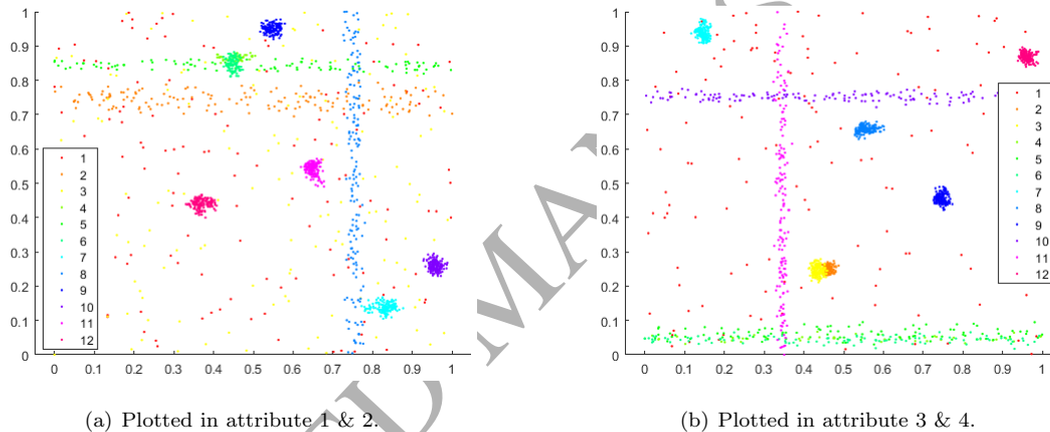


Figure 5: Distributions of the S1500 dataset in two subspaces.

to yield each attribute to be in  $[0,1]$  before the experiments began.

The parameter settings used for all clustering algorithms under comparison are shown in Table 2. They are reasonable settings as used in the literature (e.g., [8, 23, 25, 29]). For all  $k$ -means and  $k$ -medoids type clustering algorithms, we ran these algorithms over 10 trials using different random seeds for initial  $k$  centres in order to obtain their best performance.

We evaluate the clustering performance in terms of Macro F-measure, since the ground truth subspaces for each cluster are not available for real-world datasets.<sup>6</sup> Given a clustering result, we

<sup>6</sup>External evaluation measures such as Purity and NMI [2] only take into account the points assigned to clusters. A clustering, which assigns the majority of the points to noise, can produce a high score in terms of purity or NMI. F-measure is more suitable than purity and NMI in assessing clustering performance of clustering algorithms which identify noise, as in the case for subspace clustering algorithms.

Table 2: Parameters and assigned value or search range. Note that  $|C|$  is the true number of clusters.

Algorithm	Parameters and assigned value or search range
<i>CSSub</i>	$k =  C $
<i>CSSub<sub>I</sub></i>	$k =  C $ ; $\psi = 256$ ; $t = 100$
PROCLUS	$k =  C $ ; $avgDim = 0.5 \times d$
P3C	$\alpha = 0.001$ ; $poisson \in \{5, 10, \dots, 100\}$
EWKM	$k =  C $ ; $\gamma = 10$
LAC	$k =  C $ ; $h = 10$
ESSC	$k =  C $ ; $\gamma = 10$ ; $\eta = 0.5$ ; $m = s/(s - 2)$ , where $s = \min(n, d)$
FSC	$k =  C $ ; $\tau = 10$ ; $\epsilon_0 = 10^{-5}$
FG- <i>k</i> -means	$k =  C $ ; $\eta = 10$ ; $\lambda = 10$
<i>k</i> -medoids	$k =  C $

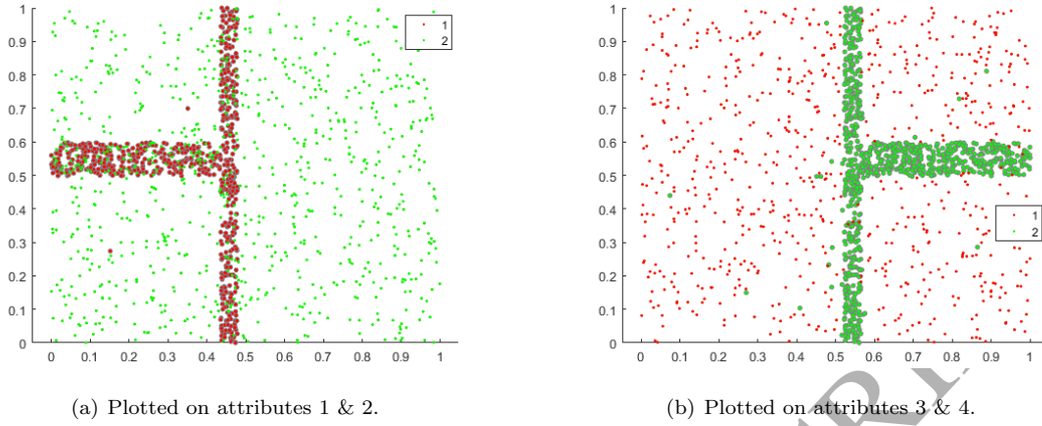
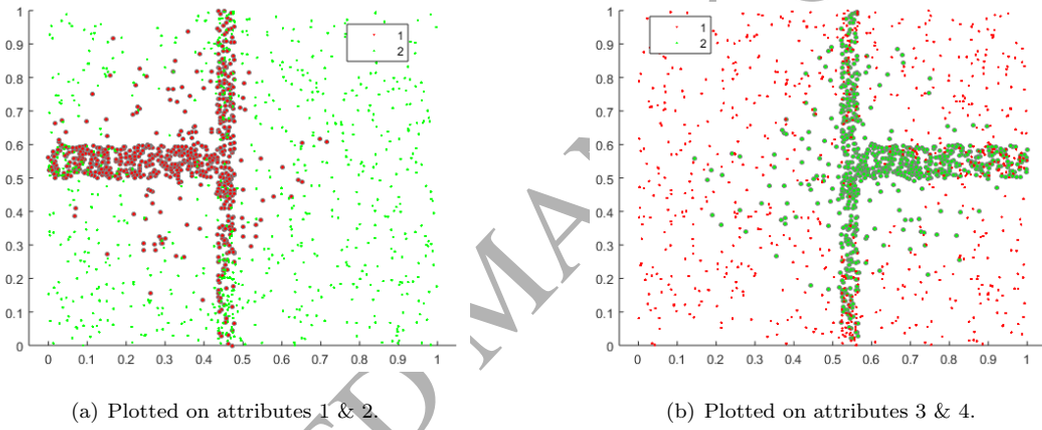
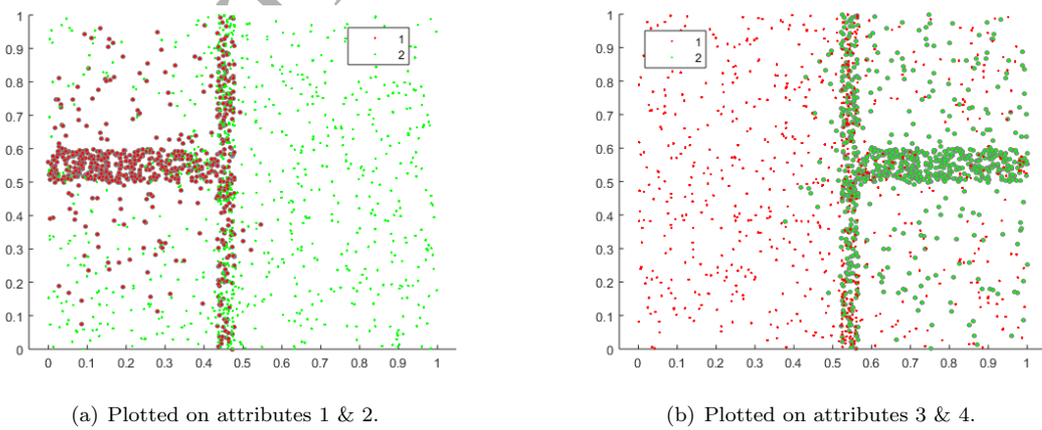
calculate the precision score  $P_i$  and the recall score  $R_i$  for each cluster  $C_i$  based on the confusion  
 335 matrix with the Hungarian method [36] for finding the maximum weight matching in a bipartite  
 graph. The F-measure of  $C_i$  is the harmonic mean of  $P_i$  and  $R_i$ . The overall F-measure is the average  
 over all clusters: F-measure =  $\frac{1}{k} \sum_{i=1}^k \frac{2P_i R_i}{P_i + R_i}$ .

### 6.1. Illustrative Clustering Performance Using a Synthetic Dataset

In this section, the 2T synthetic dataset is used to examine whether a subspace clustering algorithm  
 340 can detect arbitrarily shaped clusters in subspaces of different dimensions.

Figure 6 to Figure 8 show clusters detected by the three hard subspace algorithms on the 2T dataset.  
 Although all algorithms correctly identified the subspace for each cluster on 2T, the number of correctly  
 assigned points differs greatly among these algorithms. *CSSub* incorrectly assigned a few points only  
 (see Figure 6). Figure 7 shows that the clusters detected by PROCLUS have globular shapes—this  
 345 is the known weakness of *k*-medoids and *k*-means type algorithms when clustering algorithms are  
 performed on the given feature space—they are unable to detect arbitrarily shaped clusters. Because  
 P3C utilises a bottom-up based subspace search strategy, it may incorrectly group neighbouring points  
 which only have high densities in low dimensional projections, as shown in Figure 8. The overall result  
 is reflected in F-measure; *CSSub* has 0.94, which is much higher than those obtained by PROCLUS  
 350 (0.79) and P3C (0.76).

The relative results for the other two synthetic datasets are similar, i.e., *CSSub* has better F-  
 measure than all other algorithms. These results are summarised in the first row of Table 3. The  
 superior clustering performances of *CSSub* in detecting clusters in subspaces are mainly due to the

Figure 6: Clusters labelled by *CSSub* on the 2T dataset.Figure 7: Clusters labelled by *PROCLUS* on the 2T dataset.Figure 8: Clusters labelled by *P3C* on the 2T dataset.

use of clustering by shared subspaces—the distinctive feature of *CSSub* in comparison with other  
 355 subspace clustering algorithms. Despite the fact that the same  $k$ -medoids algorithm is used in *CSSub*,  
 the use of clustering by shared subspaces has enabled it to detect arbitrarily shaped clusters; while it  
 is impossible when  $k$ -medoids clustering is performed on attribute values of the given dataset.

## 6.2. Clustering Performance on All Datasets

In this section, we present the clustering results on all synthetic and real-world datasets. Table 3  
 360 shows the best F-measure on 15 datasets.

As we do not know whether the real-world datasets have clusters in subspaces, we perform a sanity  
 check: sort the results based on the full-space clustering  $k$ -medoids algorithm—the datasets in which  
 it has high F-measure are unlikely to have clusters in subspaces. The real world datasets in Table 3  
 are split into two subsets: one above F-measure = 0.90 and one below.

Table 3: Best F-measure of different clustering algorithms on 15 datasets; the result of real-world datasets are sorted by  
 F-measure of  $k$ -medoids. The best two performers on each dataset are boldface. The last 5 datasets are greyed because  
 full-space clustering  $k$ -medoids have performed well, indicating that they have no clusters in subspaces. *CSSub<sub>I</sub>* is  
*CSSub* with a linear time scoring function, discussed in Section 6.4. The first 3 datasets are synthetic datasets and the  
 rest are real-world datasets. The average F-measure is over the first 10 datasets.

Dataset	Hard subspace clustering				Soft subspace clustering					Full $k$ -medoids
	<i>CSSub</i>	<i>CSSub<sub>I</sub></i>	PROCLUS	P3C	EWKM	LAC	ESSC	FSC	FG- $k$ -means	
2T	<b>0.94</b>	<b>0.92</b>	0.79	0.76	0.73	0.64	0.78	0.68	0.78	0.69
S1500	<b>0.96</b>	<b>0.89</b>	0.75	0.66	0.65	0.69	0.75	0.77	0.70	0.68
D50	<b>0.89</b>	<b>0.84</b>	0.67	0.59	0.65	0.78	0.72	0.81	0.65	0.62
Glass	<b>0.46</b>	0.43	0.40	0.28	0.40	0.29	<b>0.44</b>	0.33	0.37	0.33
Liver	0.49	0.47	<b>0.54</b>	0.36	0.43	0.46	0.44	0.44	0.43	<b>0.50</b>
ILPD	<b>0.59</b>	0.44	0.56	<b>0.58</b>	0.56	0.56	0.57	0.57	0.56	0.51
Musk	0.49	0.53	0.51	0.36	0.55	0.53	0.55	<b>0.56</b>	<b>0.56</b>	0.54
Sonar	0.55	0.57	<b>0.60</b>	0.46	0.49	0.52	0.56	<b>0.61</b>	0.51	0.55
Ionosphere	<b>0.78</b>	<b>0.75</b>	0.73	0.28	0.70	0.70	0.70	0.68	0.57	0.70
Spam	0.76	0.75	0.69	0.38	0.63	0.37	<b>0.81</b>	<b>0.85</b>	0.42	0.73
Iris	0.40	0.35	0.86	0.56	<b>0.90</b>	0.88	0.90	0.96	0.89	<b>0.90</b>
Wine	0.71	0.57	0.88	<b>0.93</b>	0.95	0.95	0.95	0.94	<b>0.92</b>	0.91
Dermatology	0.70	0.73	<b>0.86</b>	0.56	0.67	0.52	0.96	0.55	0.46	<b>0.92</b>
WDBC	0.87	0.73	0.84	0.59	0.92	0.92	0.92	<b>0.94</b>	0.92	<b>0.93</b>
Breast	0.80	0.76	0.90	0.90	0.95	0.95	<b>0.95</b>	0.95	<b>0.95</b>	0.95
<i>Average</i>	0.69	0.66	0.62	0.47	0.58	0.55	0.63	0.63	0.55	0.59

365 It is indeed the case that few of the subspace clustering algorithms can perform better than  $k$ -  
 medoids full-space clustering in datasets having F-measure  $> 0.90$ . We can safely assume that the  
 last 5 datasets in Table 3 have no clusters in subspaces. We thus focus our comparison hereafter on  
 the first 10 datasets in which clusters may exist in subspaces to examine the capability of subspace  
 clustering algorithms.

370 Note that most hard subspace clustering algorithms assume that clusters exist in different subspaces. If all clusters can be separated in full space only, then they may perform badly because some important attribute values may be lost in (dis)similarity calculation. We have observed this phenomenon on some real-world datasets.

On the 10 datasets, *CSSub* has the highest average F-measure of 0.69, as shown in Table 3. Both 375 *ESSC* and *FSC* have the average F-measure of 0.63. The average F-measure of *PROCLUS* and *P3C* are 0.62 and 0.47, respectively. It is interesting to note that the average F-measure of *k*-medoids is 0.59, which is slightly higher than that of *EWKM*, *LAC* and *FG-k-means*.

We conduct the Friedman test with the post-hoc Nemenyi test [37] to examine whether the difference between any two subspace clustering algorithms is significant in terms of their average ranks. Figure 9 380 shows five subspace clustering algorithms' average ranks and critical difference. This test shows that *CSSub*, *CSSub<sub>l</sub>* and *PROCLUS* are significantly better than *P3C*. *CSSub* is significantly better than *FG-k-means* as well. *ESSC* is the best performer of soft subspace clustering algorithms, followed by *FSC*.

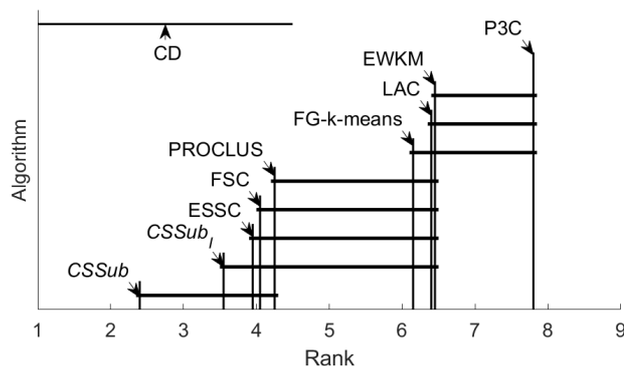


Figure 9: Critical difference (CD) diagram of the post-hoc Nemenyi test ( $\alpha = 0.10$ ). The difference between two algorithms is significant if the gap between their ranks is larger than CD. There is a line between two algorithms if the rank gap between them is smaller than CD.

385 Generally, hard subspace clustering algorithms usually perform poorly or even cannot run in a reasonable time on high-dimensional datasets (thousands of dimensions)<sup>7</sup>. *CSSub* can perform well on low-dimensional to medium-dimensional datasets (tens of dimensions) with  $n \gg d$ . This is because it examines a large proportion of all possible subspaces to identify clusters. In addition, density estimation is more reliable in low-dimensional spaces, while it becomes difficult in high-dimensional

<sup>7</sup>Although many existing hard subspace clustering algorithms are claimed to be able to deal with “high-dimensional” datasets. This usually refers to tens of dimensions only [1, 8, 38]. In this paper, we refer these “high-dimensional” datasets as medium-dimensional datasets.

spaces since these spaces are mostly empty [39].

390 Different from hard subspace clustering algorithms, many soft subspace clustering algorithms are designed to cluster high-dimensional datasets since they usually have the linear time complexity in both data size and dimensionality [23]. The examples of experimental results on real-world high-dimensional datasets are given in the Appendix B.

### 6.3. Scalability

395 We test the scalability of the subspace clustering algorithms on a synthetic dataset with increasing data size. For a fair evaluation, we set their parameters to explore the same number of subspaces, i.e., the maximum search depth is equal to the number of dimensions of the dataset used. Note that since all the five soft subspace clustering algorithms have the linear time complexity to the data size, we only report the evaluation results of ESSC as the soft subspace clustering representative.

400 Table 4 presents the runtime of the scalability test. The runtime ratio of this test is shown in Figure 10. For the increasing data size test, *CSSub*, PROCLUS and P3C have the trend of quadratic time whereas ESSC has a linear time trend. *CSSub* has the worst runtime due to the searching of best values  $\epsilon_s$  for density estimation in each subspace. However, we show in the next section that its runtime can be significantly improved by simply replacing the quadratic time density score with a  
405 linear time scoring function.

Table 4: Runtime (in seconds) of the scalability test wrt the number of points on a synthetic dataset with 4 attributes. *CSSub*, *CSSub<sub>l</sub>* and ESSC were tested in Matlab, while PROCLUS and P3C were tested in Weka.

Data size	<i>CSSub</i>	<i>CSSub<sub>l</sub></i>	PROCLUS	ESSC	P3C
500	0.62	0.59	0.01	0.01	0.03
1500	4.76	0.68	0.03	0.01	0.21
2500	12.90	0.84	0.06	0.02	0.48
3500	25.11	0.97	0.10	0.03	0.92
4500	41.48	1.08	0.15	0.03	1.51
5500	62.11	1.33	0.29	0.04	2.35

### 6.4. *CSSub* with a Linear Time Scoring Function

*CSSub* is flexible enough to employ various methods for defining clusters. It can use different scoring functions to create different types of clusters.

410 In addition, although the overall time complexity of *CSSub* is quadratic to data size, the total runtime of *CSSub* is dominated by the subspace assessment. The flexibility to use a different scoring function enables *CSSub* to easily convert from one with quadratic time complexity to one with linear time complexity by using a linear scoring function.

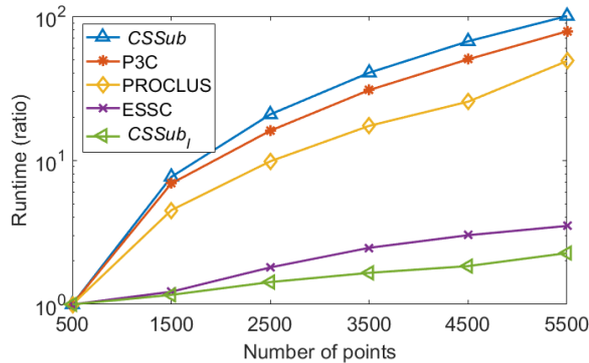


Figure 10: Runtime ratio of Scalability test.

Instead of using density score,  $CSSub_I$  uses a linear scoring function based on isolation path length score<sup>8</sup>. This score was previously used to assess subspaces for outlying aspect mining [32].

415 An *iForest* (consists of  $t$  *iTrees*) is generated for each subspace. Each *iTrees* (isolation tree) is a binary tree and is built independently using a subset  $\mathcal{D}$ , sampled without replacement from  $\pi_s(D)$ , where  $|\mathcal{D}| = \psi$ . The idea is to isolate each point from the rest of the points in the sample. This is done in a random process. At each internal node of a tree, it generates a binary splitting by random selecting an attribute and its splitting point, and then splits the sample into the two children nodes.  
420 The process is repeated recursively until no further splits can be made.

The path length of a query  $x$  on an *iTree* is the number of nodes  $x$  traverses from the root node to a leaf node. The final isolation path length score  $L_s(x)$  of  $x$  in a subspace  $s \in S$  is the average path length over  $t$  *iTrees*. We set  $\psi=256$  and  $t=100$  as the default parameter for building *iForest*, as suggested by Liu et al [40]. The details of the *iTree* building process and path length score calculation  
425 are provided in the Appendix A.

The advantages of using isolation path length as scoring function are (i) it is dimensionality unbiased [32]; and (ii) it can be computed more efficiently than density estimation—the time complexity for scoring all points in a subspace is  $O(nt \log(\psi))$ .

We found that  $CSSub_I$  is a promising method which improves  $CSSub$ 's runtime significantly  
430 and exhibits linear time behaviour wrt data size, as shown in Figure 10. In addition,  $CSSub_I$  has

<sup>8</sup>This score is derived from *iForest* (isolation forest) [40], originally used for anomaly detection as the anomaly score. *iForest* identifies anomalies as points (located in sparse regions) having short average path lengths, while normal points (located in dense regions) having long average path lengths. Because short path lengths indicate anomalies and long path lengths indicate normal points, a cluster can then be defined using neighbouring points which have path lengths longer than a certain threshold.

clustering accuracy with the average F-measure of 0.66 on the 10 datasets, which is only slightly worse than *CSSub* with F-measure of 0.69, as shown in Table 3.

### 6.5. Robustness Against Noise

*CSSub* is robust against noise because it uses an adaptive density threshold for core point identification in each subspace. The noise points were not selected by *CSSub* as core points in most subspaces. As a result, the noise points have little or no effect on its clustering result. In terms of noise attributes, *CSSub* can still perform well when only a small number of shared subspaces contain noise attributes, because the dissimilarity calculation is based on many shared subspaces. We examined this capability by adding noise points and noise attributes (separately) to the 2T dataset, and found that *CSSub* can tolerate the same level of noise as least as good as other subspace clustering algorithms, as shown in Figure 11.

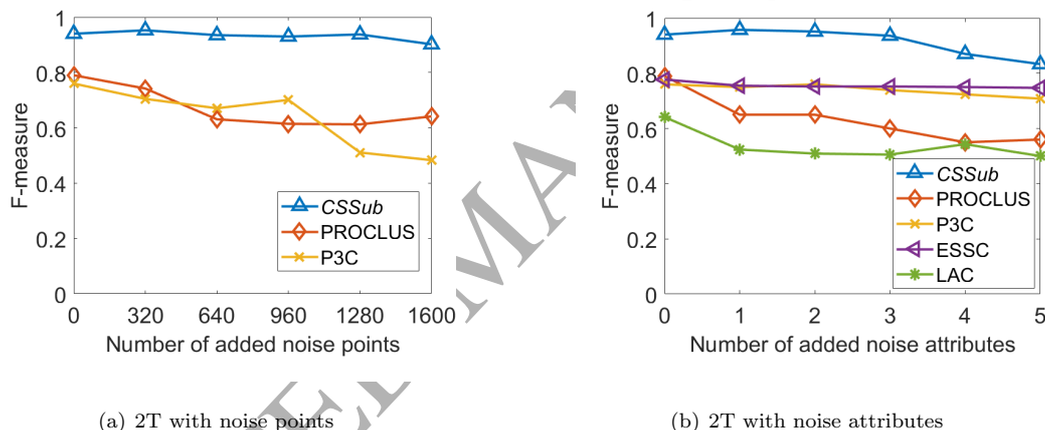


Figure 11: Best F-measure on the 2T dataset with added noise points and noise attributes. A noise point or noise attribute has values uniformly distributed in  $[0,1]$  in each attribute. Soft subspace clustering are omitted in (a) as they are very sensitive to noise points, similar to  $k$ -means.  $FG$ - $k$ -means,  $EWKM$  and  $FSC$  are omitted in (b) as they perform very similar to  $ESSC$ .

## 7. Discussion

*CSSub* is effective on low-to-medium dimensional datasets having subspace clusters with different irrelevant attributes. The experiment using the three artificial datasets, 2T, S1500 and D50, indicates that *CSSub* is particularly good at identifying subspace clusters of non-globular shape. The robustness analysis shows that it is tolerant to both noise points and noise attributes.

The *CSSub* framework provides a flexible platform for developing new subspace clustering algorithms which group points by shared subspaces. It allows different combinations of components from

either new or existing clustering algorithms. In the following, we discuss additional features of *CSSub*,  
 450 potential improvements and research directions based on this framework.

### 7.1. Subspace generation methods

*CSSub* has the flexibility of selecting systematic or non-systematic ways to generate subspaces in Stage 1. In the experiments reported thus far, the set of subspaces was generated by exhaustive enumeration to the largest subspace dimensionality  $d_{max}$  (which is set automatically). Since this  
 455 subspace generation process has the time complexity of  $O(d^{d_{max}})$ , it is infeasible on datasets with large dimensionality.

An alternative is to use a beam search method [41] for subspace generation. We design a heuristic search procedure using the beam search with the beam width  $W$  and the stage level  $l$ , described as follows. In the first level, all 1-dimensional subspaces are assigned scores and then only the top  $W$   
 460 subspaces with the highest scores are kept for the next level. Here, the score for each subspace is the maximum variance of the normalised densities, obtained from the same adaptive  $\epsilon$ -density estimator as discussed in Section 4. In the subsequent level  $(l+1)$ , we combine the  $W$  subspaces from level  $l$  in a pairwise fashion, producing new subspaces containing the union of the attributes present in the two constituent subspaces from the previous iteration. This process generates up to  $\binom{W}{2}$  new subspaces.  
 465 Each subspace is assigned scores and ranked; and only the top  $W$  subspaces with the highest scores are kept for the next level. This procedure continues until a user-defined maximum level  $H$  is reached. The  $W$  top ranked subspaces from each level  $l = \{1, \dots, H\}$  are then combined to form the set of candidate subspaces (of maximum size  $HW$ ) for use in clustering. The time complexity of this beam search method is  $O(d + HW^2)$ . Note that the proposed use of the beam search method is to generate  
 470 subspaces only in Stage 1, the other stages remain the same for *CSSub*.

Figure 12 shows the *CSSub* clustering results using the above beam search for subspace generation on the D50 dataset. We set the maximum level  $H = 10$ , and vary the beam width  $W$  from 5 to 50. It can be seen from Figure 12 that both F-measure and maximum subspace dimension increase as  $W$  increases to 30.

475 To compare the performance of *CSSub* using the beam search and *CSSub* using the exhaustive enumeration for subspace generation, we tested them on the D50 dataset with different numbers of added noise attributes, as shown in Figure 13. We set  $W = 30$  and  $H = 10$  for the beam search method, and manually set  $d_{max} = 2$  for the enumeration method. It is clear that both methods are tolerant to noise attributes. The key difference is that *CSSub* using the beam search is linear to data  
 480 dimensionality; but the one using the enumeration is exponential to dimensionality, i.e., the runtime is increased by a factor of 4 when the dimensionality is doubled from 50 to 100 (having additional 50 irrelevant attributes).

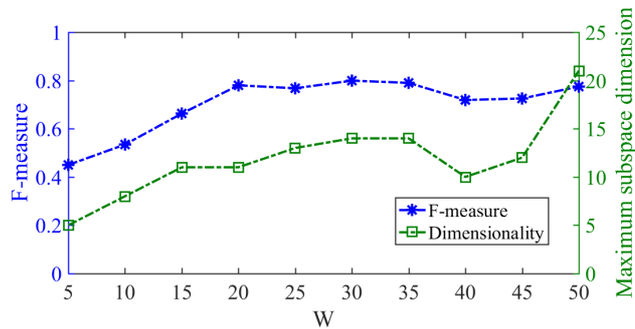


Figure 12: *CSSub* clustering results with a beam search for subspace generation on D50 dataset.

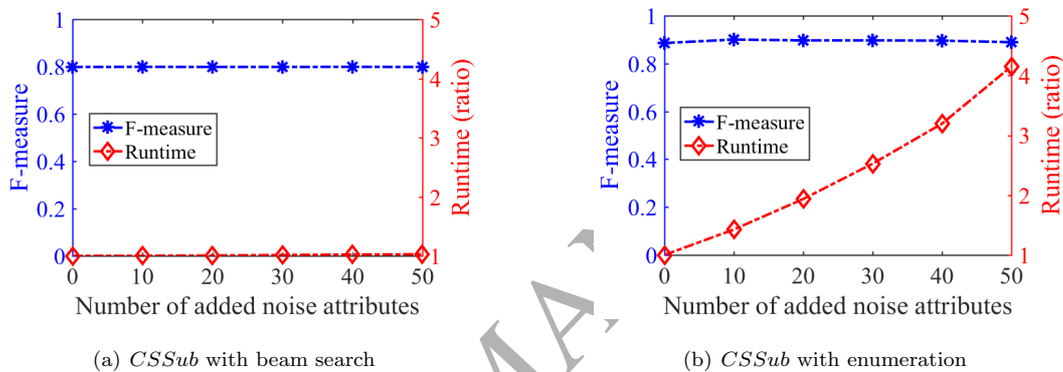


Figure 13: F-measure and runtime on the D50 dataset with added noise attributes. Each noise attribute has values uniformly distributed in  $[0,1]$ . (a) *CSSub* using the beam search for subspace generation, (b) *CSSub* using the enumeration method for subspace generation.

The beam search method has the advantage of significantly shorter runtime, especially in high dimensionality. But, it has two parameters and may lead to poor clustering results because it may lose important low-dimensional subspaces at low levels of the search. Therefore, we only recommend this method on medium-dimensional datasets.

## 7.2. Other unique features of *CSSub*

There are two other unique features of *CSSub* which have not been mentioned thus far.

First, a recent work [32] has shown that it is inappropriate to use a density score function to assess subspaces in an algorithm which employs systematic search for subspaces because a density score is a dimensionality biased function which biases toward low dimensional spaces. Existing subspace clustering methods that employ systematic search should use a criterion other than density to assess subspaces, even though they use a density-based clustering algorithm to perform clustering. For example, ENCLUS employs an entropy-based method for subspace assessment while using density-based clustering. In contrast, *CSSub* enables a density score to assess subspaces because the threshold

(i.e., mean of each subspace) employed is equivalent to a relative score which determines core points based on the mean value in each subspace, not an absolute density threshold across all subspaces. Therefore, *CSSub* has the ability to detect clusters with varied densities in different subspaces.

Second, the use of  $k$ -medoids or  $k$ -means in existing subspace clustering methods restricts the output to globular-shaped clusters only (e.g., PROCLUS and soft subspace clustering). Yet, *CSSub* has enabled  $k$ -medoids to be used to produce arbitrarily shaped clusters. This is possible because the clustering is performed on points defined by all subspaces under investigation, rather than the given feature space. In addition,  $k$ -medoids and  $k$ -means algorithms are unable to identify noise points. Yet, we show in Section 6.5 that *CSSub* using  $k$ -medoids is able to identify noise points.

### 7.3. Other Possible Refinements

We assume that only one cluster exists in each of the  $k$  subspaces detected. To examine whether the core points detected by  $k$ -medoids in a subspace have multiple clusters, density-based methods such as DBSCAN [19] can be used to separate clusters in the same subspace in the refinement stage.

It is worth mentioning that, we set  $\tau_s$  in *CSSub* as the default value for every dataset, i.e., the average score value in each subspace. We found that the clustering performance on some datasets can be significantly improved when a different  $\tau_s$  value is used.

The *CSSub* framework can be used to produce either non-overlapping clusters or overlapping clusters. One possible way to produce overlapping clusters is to extract multiple subspaces from the output of the  $k$ -medoids algorithm which covers most points of each cluster, rather than just the one which covers the maximum number of points in each cluster.

## 8. Conclusions

Grouping points by shared subspaces is a unique clustering approach which has never been attempted before, as far as we know. The approach exhibits three unique features:

1. The clustering is performed by measuring the similarity between points based on the number of subspaces they share, without examining the attribute values in the given dataset. This enables the subspace clustering to be conducted only once to produce non-redundant/non-overlapping subspace clusters. In contrast, existing algorithms produce many redundant subspace clusters from the repeated execution of clustering in a large number of subspaces because the candidate subspace selection and the clustering are tightly-coupled.
2. The approach decouples candidate subspace selection and clustering into two independent processes. As a result, it eliminates the need to have repeated runs of clustering in each of the many subspaces. The subspace selection process examines all subspaces within the limit specified for

the search, without need for a systematic search such as *Apriori*. In contrast, existing subspace clustering algorithms which have tightly-coupled processes must rely on an anti-monotonicity property to prune the search space.

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3. The decoupling approach has an added advantage that it allows different types of cluster definitions to be employed easily. The time cost of the entire process is dominated by the subspace scoring function. We show that by changing the cluster definition, which has a linear time scoring function, enables the runtime of *CSSub* to be reduced from quadratic time to linear time. A similar change is difficult, if not impossible, for existing algorithms because of the tightly coupled processes. No existing subspace clustering algorithms have the ability to use any scoring function easily, without some procedural modifications.

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The use of  $k$ -medoids or  $k$ -means in existing subspace clustering methods restricts the output to globular-shaped clusters only (e.g., PROCLUS and FG- $k$ -means). Yet, *CSSub* has enabled  $k$ -medoids to be used to produce arbitrarily shaped clusters in noisy data. This is possible because the clustering is performed on points defined by shared subspaces, rather than the given feature space.

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We show that *CSSub* discovers subspace clusters with arbitrary shapes and that it also detects noise. *CSSub* produces the best subspace clustering results on synthetic and real-world datasets in comparison with three state-of-the-art non-redundant subspace clustering algorithms which employ systematic search and optimisation for subspace search.

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Like existing hard subspace clustering algorithms, *CSSub* is meant for low-to-medium dimensionality only. How to extend *CSSub* to deal with high-dimensional datasets is an interesting future work.

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## Appendix A. Algorithm to produce isolation path length

An *iForest* consists of  $t$  *iTrees*, and each *iTree* is built independently using a subset  $\mathcal{D}$ , sampled without replacement from  $\pi_s(\mathbb{D})$ , where  $|\mathcal{D}| = \psi$ . The maximum tree height is set to  $h = \lceil \log_2 \psi \rceil$ . Note that the parameter  $e$  in *iTree* is initialised to 0 at the beginning of the tree building process.

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**Algorithm 3**  $iTree(X, e, h)$

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**Input:**  $X$  - input data;  $e$  - current height;  $h$  - height limit.

**Output:** an  $iTree$ .

```

1: if  $e \geq h$  OR  $|X| \leq 1$  then
2:   return  $exNode\{Size \leftarrow |X|\}$ ;
3: else
4:   Randomly select an attribute  $a$  from  $X$ ;
5:   Randomly select a split point  $b$  between  $min$  and  $max$  values of attribute  $a$  in  $X$ ;
6:    $X_l \leftarrow filter(X, a < b)$ ;
7:    $X_r \leftarrow filter(X, a \geq b)$ ;
8:   return  $inNode\{Left \leftarrow iTree(X_l, e + 1, h),$ 
            $Right \leftarrow iTree(X_r, e + 1, h),$ 
            $SplitAttr \leftarrow a, SplitValue \leftarrow b\}$ ;
9: end if

```

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The procedure to grow an  $iTree$  is shown in Algorithm 3. We set  $\psi=256$  and  $t=100$  as the default parameter for building  $iForest$ , as suggested by Liu et al [40].

560 The isolation path length score  $L_s(x)$  of a point  $x$  in a subspace  $s \in S$  is measured as the average path length over each  $iTree$  as follows:

$$L_s(x) = \frac{1}{t} \sum_{i=1}^t \ell_s^i(x) \quad (\text{A.1})$$

where  $\ell_s^i(x)$  is the path length of  $x$  in tree  $i$  built in  $s$ .

The path length is the number of nodes  $x$  traverses an  $iTree$  from the root node until the traversal is terminated at a leaf node. Note that if a leaf node contains more than one points  $F \subset \mathcal{D}$ , the path length of that leaf node is adjusted by adding  $2(\ln|F| + \gamma) - 2$ , where is  $\gamma \approx 0.58$  is the Euler constant  
565 [40].

## Appendix B. Clustering on high-dimensional datasets

We selected 7 real-world high-dimensional datasets for clustering evaluation <sup>9</sup>. The properties of these datasets are presented in Table B.5.

570 Table B.6 shows the best F-measures of different clustering algorithms on 7 datasets. Generally, the soft subspace clustering algorithms perform the best on these high-dimensional datasets. LAC has

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<sup>9</sup>HumanActivity and Isolet datasets are from UCI Machine Learning Repository [35]. Other datasets are from scikit-feature repository [42].

Table B.5: The properties of 7 high-dimensional datasets.

Dataset	#Points	#Dimensions	#Clusters
HumanActivity	1492	561	6
Isolet	1560	617	26
Lung	203	3312	5
GLIOMA	50	4434	4
TOX_171	171	5748	4
Prostate_GE	102	5966	2
ALLAML	72	7129	2

the highest average F-measure of 0.60. Both ESSC and  $k$ -medoids have the average F-measure of 0.58, which is higher than other subspace clustering algorithms. The hard subspace clustering algorithms  $CSSub$ ,  $CSSub_I$  and PROCLUS, which are meant for low-to-medium dimensionality only, are the worst performers on these high-dimensional datasets. This is because they only explore a tiny fraction of the total number of subspaces for a high-dimensional dataset. Note that P3C cannot get the results in reasonable time due to its exponential time complexity in dimensionality.

Table B.6: Best F-measures of different clustering algorithms on 7 datasets. The best two performers on each dataset are boldface.

Dataset	Hard subspace clustering				Soft subspace clustering					Full
	$CSSub$	$CSSub_I$	PROCLUS	P3C	EWKM	LAC	ESSC	FSC	FG- $k$ -means	$k$ -medoids
HumanActivity	0.36	0.43	0.31	—	<b>0.32</b>	0.47	0.33	0.37	<b>0.50</b>	<b>0.55</b>
Isolet	0.27	0.37	0.42	—	0.45	<b>0.55</b>	0.47	0.14	0.45	<b>0.64</b>
Lung	0.51	0.55	0.53	—	<b>0.78</b>	0.67	<b>0.81</b>	0.57	0.67	0.63
GLIOMA	0.41	0.39	0.47	—	<b>0.67</b>	0.57	<b>0.67</b>	0.50	0.51	0.54
TOX_171	0.18	0.21	0.44	—	0.47	0.49	0.45	<b>0.49</b>	<b>0.49</b>	0.42
Prostate_GE	0.57	<b>0.59</b>	0.51	—	0.58	0.57	<b>0.58</b>	0.56	0.57	0.56
ALLAML	0.55	0.44	0.69	—	0.73	<b>0.75</b>	0.72	0.71	<b>0.75</b>	0.71
<i>Average</i>	0.41	0.42	0.48	—	0.57	0.58	0.58	0.48	0.56	0.58

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