# DATA MINING 2 Transactional Clustering

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a.a. 2024/2025



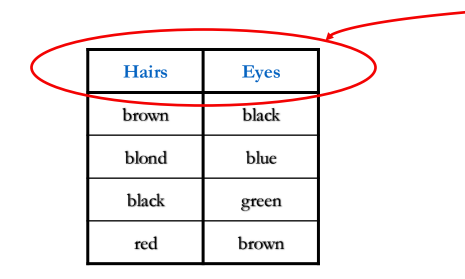
# Clustering

- **Clustering**: Grouping of objects into different sets, or more precisely, the partitioning of a data set into subsets (clusters), so that the data in each subset (ideally) share some common trait often proximity according to some defined distance measure
- Common distance functions:
  - Euclidean distance, Manhattan distance, ...
- This kind of distance functions are suitable for **numerical data**

### Not Only Numerical Data

 Acceleration 0-100 (s)	Lenght	Width	Height	Price	$\square$
0-100 (8)	(m)	<b>(</b> m <b>)</b>	(m)	(€)	
12	4	1.6	1.7	20'000	
14	3.7	1.5	1.65	16'000	Numerical Data
15	3.5	1.5	1.6	12'000	
9.4	4.2	1.8	1.7	<b>24'</b> 000	

Categorical Data



### **Boolean and Categorical Attributes**

- A **boolean** attribute corresponding to a single item in a transaction, if that item appears, the boolean attribute is set to '1' or '0' otherwise.
- A categorical attribute may have several values, each value can be treated as an item and represented by a boolean attribute.

#### Market Basket Data

- A transaction represents one customer, and each transaction contains set of items purchased by the customer.
- Clustering customers reveals customers with similar buying patterns putting them into the same cluster.
- It is useful for
  - Characterizing different customer groups
  - Targeted Marketing
  - Predict buying patterns of new customers based on profile
- A market basket database: Attributes of data points are non-numeric, transaction viewed as records with boolean attributes corresponding to a single item (TRUE if transaction contain item, FALSE otherwise).
- Boolean attributes are special case of Categorical attributes.

# **Shortcomings of Traditional Clustering**

- For categorical data we:
  - Define new criterion for *neighbors* and/or *similarity*
  - Define the ordering criterion
- Consider the following 4 market basket transactions



• using Euclidean distance to measure the closeness between all pairs of points, we find that d(P1,P2) is the smallest distance: it is equal to 1

# Shortcomings of Traditional Clustering

- If we use a hierarchical algorithm then we merge P1 and P2 and get a new cluster (P12) with (1, 1, 0.5, 1) as a centroid
- Then, using Euclidean distance again, we find:
  - d(p12,p3)= √3.25
  - d(p12,p4)= √2.25
  - d(p3,p4)= √2
- So, we should merge P3 and P4 since the distance between them is the shortest.
- However, T3 and T4 don't have even a single common item.
- So, using distance metrics as similarity measure for **categorical** data is not appropriate.

P1= (1, 1, 1, 1) P2= (1, 1, 0, 1) P3= (0, 0, 1, 0) P4= (0, 0, 0, 1)

# Algorithms for Categorical/Transactional Data

- K-Modes
- ROCK
- CLOPE
- TX-Means

#### K-Modes

Minimise 
$$P(W, Q) = \sum_{l=1}^{k} \sum_{i=1}^{n} w_{i,l} d(X_i, Q_l)$$
  
subject to  $\sum_{l=1}^{k} w_{i,l} = 1, \quad 1 \le i \le n$   
 $w_{i,l} \in \{0, 1\}, \quad 1 \le i \le n, \ 1 \le l \le k$ 

- $X = \{X_1, ..., X_n\}$  is the dataset of objects.
- $X_i = [x_1, ..., x_m]$  is an object i.e., a vector of *m* categorical attributes
- W is a matrix  $n \times k$ , with  $w_{i,l}$  equal to 1 if  $X_i$  belongs to Cluster l, 0 otherwise.
- Q = { Q<sub>1</sub>,..., Q<sub>k</sub> } is the set of representative objects (mode) for the k clusters.
- $d(X_i, Q_i)$  is a distance function for objects in the data

#### K-Modes: Distance

• K-Means as distance uses Euclidean distance

$$d(X, Y) = \overset{m}{\underset{i=1}{\overset{m}{a}}} (x_i - y_i)^2$$

 K-Modes as distance uses the number of mismatches between the attributes of two objects.

$$d_1(X, Y) = \sum_{j=1}^m \delta(x_j, y_j)$$

$$\delta(x_j, y_j) = \begin{cases} 0 & (x_j = y_j) \\ 1 & (x_j \neq y_j) \end{cases}$$

#### K-Modes: Mode

- K-Modes uses the mode as representative object of a cluster
- Given the set of objects in the cluster C<sub>1</sub> the mode is get computing the max frequency for each attribute

$$f_r(A_j = c_{l,j} | X_l) = \frac{n_{c_{l,k}}}{n}$$

### K-Modes: Algorithm

- 1. Randomly select the initial objects as modes
- 2. Scan of the data to assign each object to the closer cluster identified by the mode
- 3. Re-compute the mode of each cluster
- 4. Repeat the steps 2 and 3 until no object changes the assigned cluster

# ROCK: RObust Clustering using linK

- ROCK is a **hierarchical** algorithm for clustering transactional data (market basket databases)
- ROCK uses links to cluster instead of the classical distance notion
- ROCK uses the notion of neighborhood between pair of objects to identify the number of links between two objects

#### Input:

A set S of data points Number of *k* clusters to be found The similarity threshold

#### **Output:**

Groups of clustered data

The ROCK algorithm is divided into three major parts:

- 1. Draw a random sample from the data set
- 2. Perform a hierarchical agglomerative clustering algorithm
- 3. Label data

#### Draw a random sample from the data set:

- Sampling is used to ensure scalability to very large data sets
- The initial sample is used to form clusters, then the remaining data on dataset is assigned to these clusters

#### Perform a hierarchical agglomerative clustering algorithm:

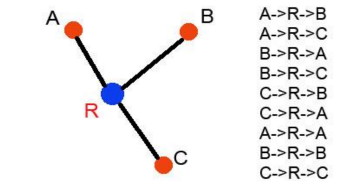
- ROCK performs the following steps which are common to all hierarchical agglomerative clustering algorithms, but with different definition to the similarity measures:
  - 1. Places each single data point into a separate cluster
  - 2. Compute the similarity measure for all pairs of clusters
  - 3. Merge the two clusters with the highest similarity (goodness measure)
  - 4. Verify a stop condition. If it is not met, then go to step 2.

### **ROCK: The Neighbors Concept**

- It captures a notion of **similarity** 
  - A and B are neighbors if sim(A, B) ≥ θ
- ROCK uses the Jaccard coefficient
  - sim(A, B)= |A ∩ B| / | A ∪ B |

A = { 1, 3, 4, 7 }  
B = { 1, 2, 4, 7, 8 }  
$$Sim(A, B) = \frac{3}{6} = \frac{1}{2} = 0.5$$

- A **link** defines the number of common neighbors between two objects:
- link(A, B) = |neighbor(A) ∩ neighbor(B) |
- Higher values of *link(A, B)* means higher probability that A and B belong to the same cluster
- **Similarity** is **local** while **link** is capturing **global** information
- A point is considered a neighbor of itself
- There is a link from each neighbor of the "root" point back to itself through the root
- Therefore, if a point has *n* neighbors, then *n*<sup>2</sup> links are due to it.



# **ROCK: Example**

- Data consisting of 6 Attributes:
  - {Book}
  - {Water, Sun, Sand, Swimming}
  - {Water, Sun, Sand, Reading}
  - {Reading, Sand}
- Resulting Jaccard Coefficient Matrix
- Set Threshold = 0.2. Neighbors:
  - N(A)={A}; N(B)={B,C,D}
  - N(C)={B,C,D}, N(D) = {B,C,D}
- Number of Links Table
  - Link (B, C) = |{B,C,D}| = 3
- Resulting Clusters after applying ROCK: {A}, {B,C,D}

{Book, Water, Sun, Sand, Swimming, Reading}

	Α	В	С	D
Α	1	0	0	0
B	0	1	0.6	0.2
С	0	0.6	1	0.5
D	0	0.2	0.5	1

	Α	В	С	D
Α	1	0	0	0
В	0	3	3	3
С	0	3	3	3
D	0	3	3	3

#### **ROCK – Criterion Function**

Maximize 
$$E_l = \sum_{i=1}^k n_i * \sum_{p_q, p_r \in C_i} \frac{link(p_q, p_r)}{n_i^{1+2f(\theta)}}$$

Dividing by the number of expected links between pairs of objects in the cluster C<sub>i</sub> we avoid that objects with a low number of links are assigned all to the same cluster

 $f(\theta) = -$ 

Where  $C_i$  denotes cluster i  $n_i$  is the number of points in  $C_i$ k is the number of clusters  $\theta$  is the similarity threshold

This goodness measure helps to identify the best pair of clusters to be merged during each step of ROCK.

$$g(C_i, C_j) = \underbrace{link[C_i, C_j]}_{(n_i + n_j)^{1+2f(\theta)} - n_i^{1+2f(\theta)} - n_j^{1+2f(\theta)} - n_j^{1+2f(\theta)}}_{\text{Number of expected cross-links between two clusters}}$$

#### Label data

- Finally, the remaining data points are assigned to the clusters.
- This is done by selecting a random sample L<sub>i</sub> from each cluster C<sub>i</sub>, then we assign each point p to the cluster for which it has the strongest linkage with L<sub>i</sub>.

#### **ROCK Summary**

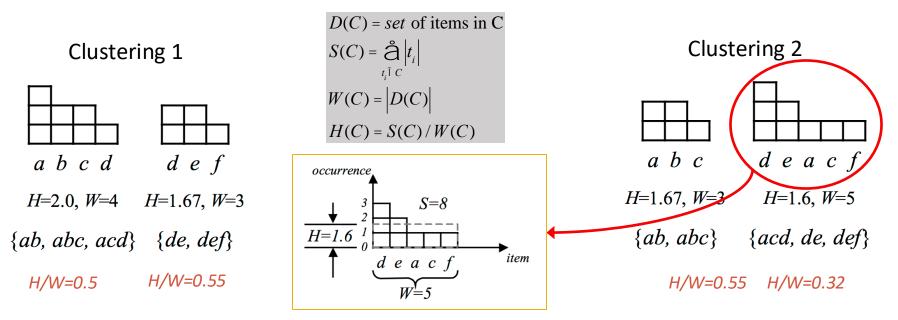
Input: dataset, number of clusters.

- 1. Draw a random sample from the data set
- 2. Places each data point into a separate cluster
- 3. Compute the similarity measure for all pairs of clusters
- 4. Merge the two clusters with the highest similarity
- 5. Verify a stop condition. If it is not met, then go to step 2.
- 6. Assign not used points to clusters using linkage similarity with respect to selected samples from each cluster

# CLOPE: Clustering with sLOPE

- Transactional clustering efficient for high dimensional data
- Uses a **global criterion function** that tries to increase the intra-cluster overlapping of transaction items by increasing the height-to-width ratio of the cluster histogram.

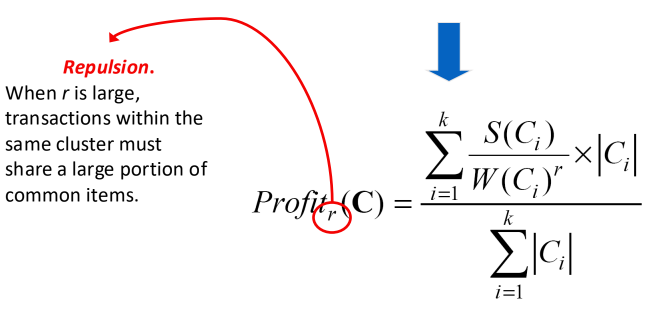
Example: 5 transactions {a,b} {a,b,c} {a,c,d} {d,e} {d,e,f}



#### Higher H/W means higher item overlapping

#### **CLOPE: Criterion Function**

- For evaluating the goodness of a clustering the gradient of a cluster is
- $G(C)=H(C)/W(C)=S(C)/W(C)^{2}$



### CLOPE: Algorithm

- /\* Phrase 1 Initialization \*/
- 1: while not end of the database file
- 2: read the next transaction  $\langle t, \text{unknown} \rangle$ ;
- 3: put t in an existing cluster or a new cluster  $C_i$  that maximize profit;
- 4: write  $\langle t, i \rangle$  back to database;
  - /\* Phrase 2 Iteration \*/
- 5: repeat
- 6: rewind the database file;
- 7: moved = false;
- 8: while not end of the database file
- 9: read  $\langle t, i \rangle$ ;
- 10: move t to an existing cluster or new cluster  $C_j$  that maximize profit;
- 11: **if**  $C_i \neq C_j$  **then**
- 12: write  $\langle t, j \rangle$ ;
- 13: *moved* = **true**;
- 14: **until** not *moved*;

### **CLOPE Summary**

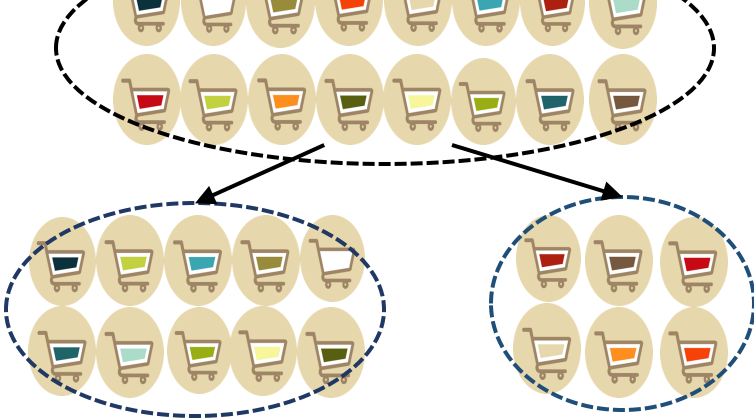
Input: dataset, repulsion, maximum number of clusters

- Phase 1
- 1. For each transaction, add it to a new cluster or to an existing one such that the profit is maximized
- Phase 2
- 1. For each transaction, try to move it to another cluster and do it if this maximizes the profit
- 2. Repeat 1. until all the transactions remain in the same cluster

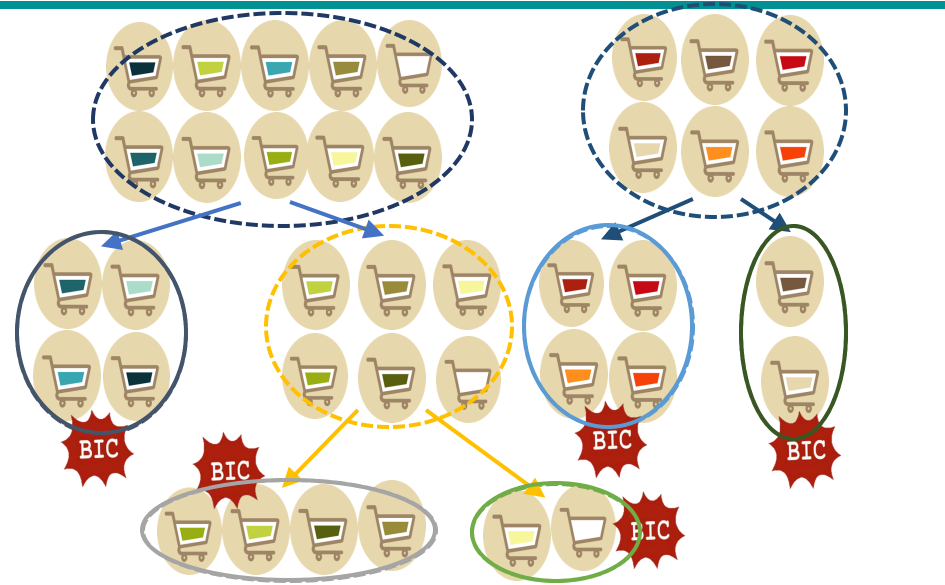


- A parameter-free clustering algorithm able to efficiently partitioning transactional data automatically
- Suitable for the case where clustering must be applied on a massive number of different datasets
  - E.g.: when a large set of users need to be analyzed individually and each of them has generated a long history of transactions
- TX-Means automatically estimates the number of clusters
- TX-Means provides the **representative transaction** of each cluster, which summarizes the pattern captured by that cluster.





#### How It Works 2/3

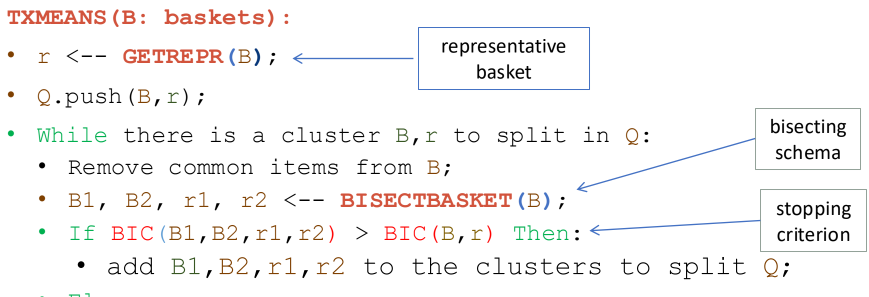


#### How It Works 3/3

• Clusters

• Representative Baskets

#### **TX-Means Algorithm**



- Else
  - add B,r to the clustering result C;
- Return C;

#### **Bisecting Schema**

#### **BISECTBASKET (B: baskets):**

- SSE <-- inf;
- r1,r2 <-- select random initial baskets in B as representative;</li>
- While True:
  - C1,C2 <-- assign baskets in B with respect to r1,r2;
  - r1 new <-- GETREPR(C1); r2 new <-- GETREPR(C2); 🔨
  - SSE\_new <-- SSE(C1,C2,r1\_new,r2\_new);
  - If SSE\_new >= SSE Then:
    - Return C1,C2,r1,r2;
  - r1,r2 <-- r1\_new,r2\_new;</pre>

overlap-based distance function: Jaccard coefficient

#### Get Representative Baskets

#### GETREPR(B: baskets):

- I <-- not common items in B;
- r <-- common items in B;</li>
- While I is not empty:
  - Add to r the items with maximum frequency in I;
  - Calculate the distance between r and the baskets in B;
  - If the distance no longer decreases Then:
    - Return r;
  - Else
    - remove from I the items with maximum frequency;
- Return r;

overlap-based distance function (Jaccard coefficient)

- Clustering of a big individual transactional dataset B.
- TX-Means is scalable thanks to the following sampling strategy.
- Sampling strategy:
  - Random selection of a subset S of the baskets in B;
  - Run of TX-Means on the subset S and obtain clusters C and representative baskets R;
  - Assign the remaining baskets B/S to the clusters C using a nearest neighbor approach with respect to the representative baskets R.

### References

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- Guidotti, R., et al. Clustering individual transactional data for masses of users. 2017.

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	Abstract	lutions for these problems. Speed is great	ly improved	A CALL AND A CALL AND
	Despite its popularity for general clustering,	by embedding the dataset in a multiresolu	tion kd-tree	The second second second second second
	K-means suffers three major shortcomings; it scales poorly computationally, the num-	and storing sufficient statistics at its node	s. A careful	CONCERNMENT OF A
	it scales poorly computationally, the num-	analysis of the centroid locations allows ric "proofs" about the Voronoi boundarie	s, and (un-	
	ber of clusters $K$ has to be supplied by the user, and the search is prone to local min-	like all of (Deng & Moore, 1995; Zhang	et al., 1995;	
	ima. We propose solutions for the first two	Moore, 1999)) there is absolutely no ap anywhere in the computation. An additio	nal geomet-	A DI LATE STOR
	problems, and a partial remedy for the third. Building on prior work for algorithmic accel	ric computation, blacklisting, maintains a	list of just	A CONTRACT OF
	eration that is not based on approximation.	those centroids that need to be considered region (Pelleg & Moore, 2000). Blacklistin	introduction to	
	we introduce a new algorithm that efficiently, searches the space of cluster locations and	extremely fast but also scales very well wi	th the num	And a state of the
	number of clusters to optimize the Bayesian Information Criterion (BIC) or the Akaike	ber of centroids, allowing tractable 10, 000- rithms. This fast algorithm is used as a bu		
	Information Criterion (BIC) or the Akaike	in X-means: a new algorithm that quick	y estimates	
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	ing cached sufficient statistics and a new very	ing local decisions about which subset of centroids should split themselves in order	a hater fa	
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	the number of classes and their parameters. Experiments show this technique reveals the		Information Systems Vol. 25, No. 5, pp. 345–3 © 2000 Elsevier Science Lid. All rights Printed in Grea 0006-4379/00 \$20.0	reserved t Britain
	true number of classes in the underlying dis-	siz ta	0306-4379/00 \$20.0	0.00+0.00
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		wi Sudipto Guha wi	<sup>1</sup> , RAJEEV RASTOGI <sup>2</sup> , and KYUSEOK SHIM <sup>3</sup>	
	K-means (Duda & Hart, 1973; Bishop, 1995) has long been the workhorse for metric data. Its attractive-	m <sup>I</sup> Stanfe	ord University, Stanford, CA 94305, USA	
<b>ר</b>	been the workhorse for metric data. Its attractive- ness lies in its simplicity, and in its local-minimum	<sup>2</sup> Bell I.	ord University, Stanford, CA 94305, USA aboratories, Murmy Hill, NJ 07974, USA	
/	convergence properties. It has, however, three main shortcomings. One, it is slow and scales poorly with	<ol> <li><sup>3</sup>Korea Advanced Institute of Science</li> </ol>	and Technology and Advanced Information Technology Research ( Taejon 305-701, Korea	Center,
<b>∠</b> .	subtroomings, one, it is now and scales poorly with			
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	fosca.giannotti@isti.cnr.it	dino.pedreschi@di.unipi.it	Transacti	onal Data
	ABSTRACT	from other users. This requires that		g Guan Jinyuan You
	Mining a large number of datasets recording human activitie	es for included in any data mining metho	Dept. of Computer Science & Engine	ering. Shanghai Jiao Tong University
	making sense of individual data is the key enabler of a new y	wave the necessity to automatically capt vidual behaviors. Due to the potent	Shanghai, 200	ering., Shanghai Jiao Tong University 030, P.R.China
	of personalized knowledge-based services. In this paper we f on the problem of clustering individual transactional data			52581638
	large mass of users. Transactional data is a very pervasive kit		{yang-yl, guan-xd, yo	ou-jy}@cs.sjtu.edu.cn
	information that is collected by several services, often invol huge pools of users. We propose txmeans, a parameter-free			
	tering algorithm able to efficiently partitioning transactional	data characteristics of the dataset under a		
	in a completely automatic way. Txmeans is designed for the	case personalized patterns from transact	ABSTRACT	The Largeltern [13] algorithm groups large categorical databases
	where clustering must be applied on a massive number of diff- datasets, for instance when a large set of users need to be anal	erent In this paper we focus on the pa tional clustering for a large number	This paper studies the problem of categorical data clustering, especially for transactional data characterized by high dimensionality and large volume. Starting from a heuristic method	by iterative optimization of a global criterion function. The criterion function is based on the notion of large item that is the
	individually and each of them has generated a long history of t	rans- collection of transactions, transacti	dimensionality and large volume. Starting from a heuristic method	item in a cluster having occurrence rates larger than a user-defined parameter minimum support. Computing the global criterion
	actions. A deep experimentation on both real and synthetic dat	asets covering groups of homogeneous t	of increasing the height-to-width ratio of the cluster histogram, we develop a novel algorithm - CLOPE, which is very fast and	function is much faster than those local criterion functions defined
	shows the practical effectiveness of txmeans for the mass clusts of different personal datasets, and suggests that txmeans ou	transactional clustering require eith	scalable, while being quite effective. We demonstrate the performance of our algorithm on two real world datasets, and	on top of pair-wise similarities. This global approach makes LargeItem very suitable for clustering large categorical databases.
	forms existing methods in terms of quality and efficiency. Fit	nally, that is not automatic, or an extrem	performance of our algorithm on two real world datasets, and compare CLOPE with the state-of-art algorithms.	In this paper, we propose a powel global criterion function that
	we present a personal cart assistant application based on txm	eans. that does not scale to large user b repeatedly applying the existing pr		tries to increase the intra-cluster overlapping of transaction items
	1 INTRODUCTION	lions of different datasets – which i	Keywords	In any paper, we propose in lover groun criterion related that tries to increase the intra-cluster overlapping of transaction items by increasing the height-to-width ratio of the cluster histogram. Moreover, we generalize the idea by introducing a parameter to
	The most disruptive effect of our always-connected society is	large population of users – is simp data, problem, i.e., the separate individua	data mining, clustering, categorical data, scalability	control the ightensio of the cluster. Different number of clusters can be obtained by varying this parameter. Experiments show that
	the digital breadcrumbs left behind us as a side effect of our ever	vday problem, i.e., the separate individua yday transactional datasets, as mass clusi		can be obtained by varying this parameter. Experiments show that
	usage of digital technologies. Thanks to these data, human activ	rities The problem to design parameter		our algorithm runs much faster than Largeltem, with clustering quality quite close to that of the ROCK algorithm [7].
	are becoming observable, measurable, quantifiable and, predic At individual level, each person generates more than 5Gb of dat	table. been addressed in the context of non a per like <i>xmeans</i> [22], which are perfe	Clustering is an important data mining technique that groups together similar data records [12, 14, 4, 1]. Recently, more	To gain some basic idea behind our algorithm, let's take a small
	year. An avalanche of information that, for the most part, cor	usists of the clustering problems. Unfort		market basket database with 5 transactions {(apple, banana), (apple, banana, cake), (apple, cake, dish), (dish, egg), (dish, egg,
	of transactions (or baskets), i.e., a special kind of categorical in the form of sets of event data, such as the items purchase	data applicable to transactional data. To	13], where records are made up of non-numerical attributes. Transactional data, like market basket data and web usage data, can be thought of a special type of categorical data having boolean	(appre, banana, case), (appre, case, dish), (asia, egg), (adam, egg), (fish), For simplicity, transaction (apple, banana) is abbreviated to ab, etc. For this small database, we want to compare the following
	in the form of sets of event data, such as the items purchase a shopping cart, the web pages visited in a browsing session		Transactional data, like market basket data and web usage data, can be thought of a special type of categorical data having boolean	ab, etc. For this small database, we want to compare the following two electering (1) ((ab abc acd)) (de doft) and (2) ((ab abc))
	songs listened in a time period, the clinical events in a pati	ent's generally not efficient and overestim	value, with all the possible items as attributes. Fast and accurate	two clustering (1) {{ab, abc, acd}, {de, def} and (2) {{ab, abc}, {acd, de, def} For each cluster, we count the occurrence of every distinct item, and then obtain the height (H) and width (W) of the
	history. Such kind of data may be key enablers of a new war knowledge-based services, and of new scientific discoveries.	ve of In addition, they do not provide repr items that characterize the transact	clustering of transactional data has many potential applications in retail industry, e-commerce intelligence, etc.	distinct item, and then obtain the height (H) and width (W) of the
	Several application contexts involve the analysis of a large	num- In this paper we propose tymean		cluster. For example, cluster $\{a, bc, acd\}$ has the occurrences of $a:3, b:2, c:2, and d:1, with H=2.0 and W=4. Figure 1 shows these$
	ber of datasets, each one characterized by different properties	For ing method providing a viable soluti	However, fast and effective clustering of transactional databases is extremely difficult because of the high dimensionality, sparsity,	results geometrically as histograms, with items sorted in reverse order of their occurrences, only for the sake of easier visual
	instance, this is the case of individual transactional data - I sales, web sessions, credit card transactions, etc where	retail a massive number of different dat	and huge volumes often characterizing these databases. Distance- based approaches like k-means [11] and CLARANS [12] are	interpretation.
	user produces historical data that need to be analyzed separ-	ately finding clusters in the specific cont-	effective for low dimensional numerical data. Their performances on high dimensional categorical data, however, are often	n n
			unsatisfactory [7]. Hierarchical clustering methods like ROCK [7]	
	classroom use is granted without fee provided that copies are not made or distr for roufit or commercial advantage and that copies have this reader and the Gar	ibuted it automatically estimates the num itation tracting the clusters, it provides th	have been demonstrated to be quite effective in categorical data	abcd def abc deacf
	on the first page. Copyrights for components of this work owned by others than much become distinguishing of the provided of the second by others than	ACM each cluster, which summarizes the	clustering, but they are naturally inefficient in processing large databases,	H=2.0, W=4 H=1.67, W=3 H=1.67, W=3 H=1.6, W=5
	to post on servers or to redistribute to lists, requires prior specific permission an	ublish, d/or a Txmeans employs a top-down divisi		{ab, abc, acd} {de, def} {ab, abc} {acd, de, def}
	Permission to make digital or hand copies of all or part of this work for perso- fasorous one is granted without for provided that copies are not made or dains on the first part, copyrights for components of this work encoursed on the first part, copyrights for components of this work encoursed to post on averses to relativative to interview of the provider works are to post on averses to relativative to interview program of the provider one first part of the provider of the provider of the provider one of the provider one first part of the provider of	a unique cluster, and then iterativ sub-clusters. Txmeans calculates th		clustering (1) clustering (2)
	© 2017 ACM. 978-1-4503-4887-4/17/08\$15.00 DOI: 10.1145/3097983.3098034	centroids of the sub-clusters by ado	Permission to make digital or hard conies of all or part of this work for	Figure 1. Histograms of the two clusterings.
			personal or classroom use is granted without fee provided that copies are	We indee the qualities of these two clusterings geometrically, by
			not made or distributed for profit or commercial advantage and that copies bear this notice and the fall citation on the first page. To copy	analyzing the heights and widths of the clusters. Leaving out the two identical histograms for cluster {de, def} and cluster {ab, abc},
			Permission to make digital or hard copies of all or part of this work for personal or classroom use is granted without fee provided that copies are not made or distributed for profix or commercial advantage and that copies bear this notice and the full citation on the first page. To copy otherwise, or republish, to part to enservers or to redistribute to lists, requires prior specific permission and/or a fee.	two identical histograms for cluster $(ae, ae)$ and cluster $(ab, abc)$ , the other two histograms are of different quality. The histogram for cluster $\{ab, abc, acd\}$ has only 4 distinct items for 8 blocks
			SIGKDD '02, July 23-26, 2002, Edmonion, Alberta, Canada.	for cluster {ab, abc, acd} has only 4 distinct items for 8 blocks (H=2.0, H/W=0.5), but the one for cluster {acd, de, def} has 5, for

X-means: Extending K-means with Efficient Estimation of the Number of Clusters

# **Exercises Transactional Clustering**

- Suppose we have four verses contains some subjects, as follows:
- P1={ judgment, faith, prayer, fair}
- P2={ fasting, faith, prayer}
- P3={ fair, fasting, faith}
- P4={ fasting, prayer, pilgrimage}
- the similarity threshold = 0.3, and number of required cluster is 2.

Using Jaccard coefficient as a similarity measure, we obtain the following similarity table

	P1	P2	P3	P4
P1	1	0.4	0.4	0.17
P2		1	0.5	0.5
<b>P</b> 3			1	0.2
P4				1

- Since we have a similarity threshold equal to 0.3, then we derive the adjacency table: →
- By multiplying the adjacency table with itself, we derive the following table which shows the number of links (or common neighbors): →

	P1	P2	P3	P4
P1	1	0.4	0.4	0.17
P2		1	0.5	0.5
P3			1	0.2
P4				1

	P1	P2	P3	P4
P1	1	1	1	0
P2		1	1	1
P3			1	0
P4				1

	P1	P2	P3	P4
P1	-	3	3	1
P2		-	3	2
P3			-	1
P4				-

 we compute the goodness measure for all adjacent points ,assuming that

• 
$$f(\theta) = 1 - \theta / 1 + \theta = 1 - 0.3 / 1 + 0.3 = 0.54$$

- $\bullet$  we obtain the following table  $\rightarrow$
- we have an equal goodness measure for merging ((P1,P2), (P2,P3), (P3,P1))

$$g(P_i, P_j) = \frac{link[P_i, P_j]}{(n+m)^{1+2f(\theta)} - n^{1+2f(\theta)} - m^{1+2f(\theta)}}$$

Pair	Goodness measure
P1,P2	1.35
P1,P3	1.35
P1,P4	0.45
P2,P3	1.35
P2,P4	0.90
P3,P4	0.45

- Now, we start the hierarchical algorithm by merging, say P1 and P2.
- A new cluster (let's call it C(P1,P2)) is formed.
- It should be noted that for some other hierarchical clustering techniques, we will not start the clustering process by merging P1 and P2, since Sim(P1,P2) = 0.4, which is not the highest. But, ROCK uses the number of links as the similarity measure rather than distance.

- Now, after merging P1 and P2, we have only three clusters. The following table shows the number of common neighbors for these clusters:→
- Then we can obtain the following goodness measures for all adjacent clusters:→

	C(P1,P2)	P3	P4
C(P1,P2)	_	3+3	2+1
P3		-	1
P4			-

Pair	Goodness measure
C(P1,P2),P3	1.31
C(P1,P2),P4	0.66
P3,P4	0.45

 Since the number of required clusters is 2, then we finish the clustering algorithm by merging C(P1,P2) and P3, obtaining a new cluster C(P1,P2,P3) which contains {P1,P2,P3} leaving P4 alone in a separate cluster.

 Given the following similarity matrix find the clustering result knowing that the similarity threshold = 0.4, and number of required cluster is 2.

	p1	p2	р3	р4	р5
p1	1	0.7	0.2	0.5	0.5
p2		1	0.6	0.8	0.1
р3			1	0.5	0.4
p4				1	0.3
p5					1

	p1	p2	р3	р4	р5
p1	1	0.7	0.2	0.5	0.5
p2		1	0.6	0.8	0.1
р3			1	0.5	0.4
p4				1	0.3
р5					1

	p1	p2	р3	р4	р5
p1	1	1	0	1	1
p2	1	1	1	1	0
р3	0	1	1	1	1
p4	1	1	1	1	0
р5	1	0	1	0	1

	p1	p2	р3	р4	р5
p1	1	1	0	1	1
p2	1	1	1	1	0
р3	0	1	1	1	1
p4	1	1	1	1	0
p5	1	0	1	0	1

	p1	p2	р3	р4	р5
p1	-	3	3	3	2
p2		-	3	4	2
р3			-	3	2
p4				-	2
р5					-

• 
$$f(\theta) = 1 - \theta / 1 + \theta = 1 - 0.4 / 1 + 0.4 = 0.43$$

•  $1 + 2 f(\theta) = 1.86$ 

$$g(P_i, P_j) = \frac{link[P_i, P_j]}{(n+m)^{1+2f(\theta)} - n^{1+2f(\theta)} - m^{1+2f(\theta)}}$$

	p1	p2	р3	р4	р5
p1	-	3	3	3	2
p2		-	3	4	2
р3			-	3	2
p4				-	2
р5					-

	p1	p2	р3	p4	р5
p1	-	1.84	1.84	1.84	1.22
p2		-	1.84	2.45	1.22
р3			-	1.84	1.22
p4				-	1.84
p5					-

• 
$$f(\theta) = 1 - \theta / 1 + \theta = 1 - 0.4 / 1 + 0.4 = 0.43$$

•  $1 + 2 f(\theta) = 1.86$ 

$$g(P_i, P_j) = \frac{link[P_i, P_j]}{(n+m)^{1+2f(\theta)} - n^{1+2f(\theta)} - m^{1+2f(\theta)}}$$

	p1	p2	р3	p4	р5		p1	p2p4	р3	р5		p1	p2p4	р3	р5
p1	-	3	3	3	2	p1	_	6	3	2	p1	_	1.94	1.84	1.22
p2		-	3	4	2	p2p4		_	6	4	p2p4			1.94	1.29
p3			-	3	2				0				_	1.94	
p4				-	2	p3			-	2	p3			-	1.22
p5					-	р5				-	р5				-

• Final Clusters: p1234 p5

# **Clope Exercise 1**

Transactions: abc, abc, ab, ad, def, ade, ade

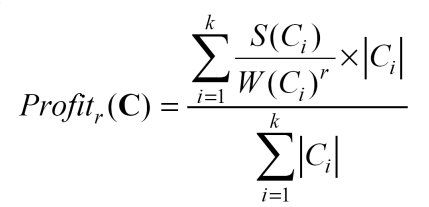
Split1:

- 4 transactions: abc, abc, ab, ad
  - a:4, b:3, c:2, d:1 -> S=10; W=4; H=10/4=2,5; H/W=2,5/4=0,625
- 3 transactions: def, ade, ade
  - a:2, d:3, e:3, f:1 -> S=9; W=4; H=9/4=2,25; H/W=2,25/4=0,56

Split2:

- 2 transactions: abc, abc, ab
  - a:3, b:3, c:2 -> S=8; W=3; H=8/3=3,6; H/W=0,88
- 2 transactions: ad, def, ade, ade
  - a:3, d:4, e:3, f:1 -> S=11; W=4; H=11/4=2,75; H/W=2,75/4=0,68

Split1 is the best clustering considering r=2 Profit(Split1) =  $(10/4^2 * 4 + 9/4^2 * 3)/7 = 0.59$ Profit(Split2) =  $(8/3^2 * 3 + 11/4^2 * 4)/7 = 0.77$ 



## **Clope Exercise 2**

Split1:

- 4 transactions: abc, abc, ab, a
  - a: 4, b:3, c: 2 -> sol: S=9; W=3; H=9/3=3; H/W=1
- 3 transactions: def, de, de
  - d: 3, e:3, f: 1 -> sol: S=7; W=3; H=7/3=2.33; H/W=0.77

Split2:

- 2 transactions: abcd, ab
  - a: 2, b:2, c: 1, d:1 -> sol: S=6; W=4; H=6/4=1.5; H/W=0.37
- 2 transactions: ec, ec
  - e:2, c: 2 -> sol: S=4; W=2; H=4/2=2; H/W=1

Split1 is the best clustering considering r=2 Profit(Split1) =  $(9/3^2 * 4 + 7/3^2 * 3)/7 = 0.90$ Profit(Split2) =  $(6/4^2 * 2 + 4/2^2 * 2)/4 = 0.16$ 

