Meta Clustering

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Abstract

Clustering is ill-defined. Unlike supervised learning where labels lead to crisp performance criteria such as accuracy and squared error, clustering quality depends on how the clusters will be used. Devising clustering criteria that capture what users need is difficult. Most clustering algorithms search for optimal clusterings based on a pre-specified clustering criterion. Our approach differs. We search for many alternate clusterings of the data, and then allow users to select the clustering(s) that best fit their needs. Meta clustering first finds a variety of clusterings and then clusters this diverse set of clusterings so that users must only examine a small number of qualitatively different clusterings. We present methods for automatically generating a diverse set of alternate clusterings, as well as methods for grouping clusterings into meta clusters. We evaluate meta clustering on four test problems and two case studies. Surprisingly, clusterings that would be of most interest to users often are not very compact clusterings.

1. Introduction

Clustering performance is difficult to evaluate [30]. In supervised learning, model performance is assessed by comparing model predictions to targets. In clustering we do not have targets and usually do not know *a priori* what groupings of the data are best. This hinders discerning when one clustering is better than another, or when one clustering algorithm outperforms another. To make matters worse, clustering often is applied early during data exploration, before users understand the data well enough to define suitable clustering criteria. This creates a chicken-or-the-egg problem where knowing how to define a good clustering criterion requires understanding the data, but clustering is one of the principal tools used to help understand the data.

This fundamental differences between supervised and unsupervised learning have profound consequences. In particular, while it makes sense to talk about the "best" model(s) in supervised learning (e.g. the most accurate

model(s)), often it does not make sense to talk about the "best" clustering. Consider a database containing information about people's age, gender, education, job history, spending patterns, debts, medical history, etc. Clustering could be applied to the database to find groups of similar people. A user who wants to find groups of consumers who will buy a car probably wants different clusters than a medical researcher looking for groups with high risk of heart disease. In exploring the same data, different users want different clusterings. No "correct" clustering exists. Moreover, theoretical work suggests that it is not possible to achieve all of the properties one might desire of clustering in a single clustering of the data [21].

Most clustering methodologies focus on finding optimal or near-optimal clusterings, according to specific clustering criteria. However, this approach often is misguided. When users cannot specify appropriate clustering criteria in advance, effort should be devoted to helping users find appropriate clustering criteria. In practice, users commonly begin by clustering their data and examining the results. They then make educated guesses about how to change the distance metrics or algorithm in order to yield a more useful clustering. Such a search is tedious and may miss interesting partitionings of the data.

In this paper we introduce *meta clustering*, a new approach to the problem of clustering. Meta clustering aims at creating a new mode of interaction between users, the clustering system, and the data. Rather than finding one optimal clustering of the data, meta clustering finds many alternate good clusterings of the data and allows the user to select which of these clusterings is most useful, exploring the space of reasonable clusterings. To prevent the user from having to evaluate too many clusterings, the many base-level clusterings are organized into a meta clustering, a clustering of clusterings that groups similar base-level clusterings together. This meta clustering makes it easier for users to evaluate the clusterings and efficiently navigate to the clustering(s) useful for their purposes.

Meta clustering consists of three steps. First, a large number of potentially useful clusterings is generated. Then a distance metric over clusterings measures the similarity between pairs of clusterings. Finally, the clusterings are themselves clustered at the meta level using the computed pairwise similarities. The clustering at the meta level allows the user to select a few representative yet qualitatively different clusterings for examination. If one of these clusterings is appropriate for the task at hand, the user may then examine other nearby clusterings in the meta level space.

An analogy may be helpful. Photoshop, the photo editing software, has a tool called "variations" that presents to the user different renditions of the picture that have different color balances, brightnesses, contrasts, and color saturations. Instead of having to know exactly what tool to use to modify the picture (which requires substantial expertise), the user only has to be able to select the variation that looks best. The selected variation then becomes the new center, and variations of it are presented to the user. This process allows users to quickly zero in on the desired image rendition. The goal in meta clustering is to provide a similar "variations" tool for clustering so that users do not have to know how to modify distance metrics and clustering algorithms to find useful clusterings of their data. Instead, meta clustering presents users with an organized set of clustering variations; users can select and then refine the variation(s) that are best suited to their purposes.

The paper proceeds as follows. Section 2 defines meta clustering. Section 2.1 describes how to generate diverse high-quality clusterings. Section 2.2 describes how to measure similarity between clusterings to cluster clusterings at the meta level. Section 3 describes four data sets used for evaluation, and Section 4 presents the empirical results. Section 5 presents a protein clustering case study. Section 6 presents a phoneme clustering case study. Section 7 covers the related work. Section 8 is a discussion and summary.

2. Meta Clustering

The approach to meta clustering presented in this paper is a sampling-based approach that searches for distance metrics that yield the clusterings most useful to the user. Algorithmic (i.e. non-stochastic) approaches to meta clustering are possible and currently are being developed.

Here we break meta clustering into three steps:

- 1. Generate many good, yet qualitatively different, base-level clusterings of the same data.
- 2. Measure the similarity between the base-level clusterings generated in the first step so that similar clusterings can be grouped together.
- Organize the base-level clusterings at a meta level (either by clustering or by low-dimension projection) and present them to users.

These steps are described in the rest of this section.

2.1. Generating Diverse Clusterings

The key insight behind meta clustering is that in many applications, data may be clustered into a variety of alternate groupings, each of which may be beneficial for a different purpose. To be useful, the alternate clusterings cannot be random partitions of the data, but must reflect *genuine structure within the data*. We follow two approaches to generate a diverse set of quality clusterings. In the first, we note that k-means generates many different reasonable clusterings (all but the "best" of which are typically discarded) because different random initializations of k-means often get stuck in different local minima. In the second approach, we apply random weights to feature vectors before clustering the data with k-means to emphasize different aspects of the data. These approaches for finding diverse clusterings are described in the remainder of Section 2.1.

2.1.1 Diverse Clusterings from K-Means Minima

K-means is an iterative refinement algorithm that attempts to minimize a squared error criterion [11]. Each cluster is initialized by setting its mean to a random point in the data set. Each step of the iterative refinement performs two tasks. First, the data points are classified as being a member of the cluster with the nearest cluster mean. Second, the cluster means are updated to be the actual mean of the data points in each cluster. This is repeated until no points change membership or for some maximum number of iterations. When no points change membership, k-means is at a local minimum in the search space: there is no longer a move that can reduce the squared error. The output of k-means is typically highly dependent on the initialization of the cluster means: the search space has many local minima [3, 5]. In practice, k-means is run many times with many different initializations, and the clustering with the smallest sum-of-squared distances between cluster means and cluster members is returned as the final result.

In meta clustering, however, we are interested in generating a wide variety of reasonable clusterings. The local minima of k-means provide a set of easily-attainable clusterings, each of which is reasonable since no point can change membership to improve the clustering. K-means can be run many times with many different random initializations, and each local minimum can be recorded. As we shall see in Section 4.2, the space of k-means local minima is small compared to the space of reasonable clusterings, so we use an additional means of generating diverse clusterings: random feature weighting.

2.1.2 Diverse Clusterings from Feature Weighting

Consider data in vector format. Each item is described by a vector of features, and each dimension in the vector is

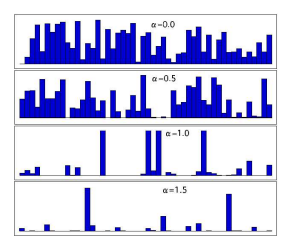


Figure 1. Zipf Distribution. Each row visualizes a Zipf distribution with a different shape parameter, α . Each row has 50 bars representing 50 random samples from the distribution, with the height of the bar proportional to the value of the sample.

a feature that will be used when calculating the similarity of points for clustering. By weighting features before distances are calculated (i.e. multiplying feature values by particular scalars), we can control the importance of each feature to clustering [34]. Clustering many times with different random feature weights allows us to find qualitatively different clusterings for the data using the same clustering algorithm.

Feature weighting requires a distribution to generate the random weights. We consider both uniform and power law distributions. Empirically, uniformly distributed weights often do not explore the weight space thoroughly. Consider the case where only a few features contain useful information, while the others are noise. It is unlikely that a uniform distribution would generate values that weight the few important variables highly while assigning low weights to the majority of the variables, while weights generated from a power law distribution can weight only a few variables highly.

We will use a Zipf power law distribution because there is empirical evidence that feature importance is Zipf-distributed in a number of real-world problems [8, 15]. A Zipf distribution describes a range of integer values from 1 to some maximum value K. The frequency of each integer is proportional to $\frac{1}{i^{\alpha}}$ where i is the integer value and α is a shape parameter. Thus, for $\alpha=0$, the Zipf distribution becomes a uniform distribution from 1 to K. As α increases, the distribution becomes more biased toward smaller numbers, with only the occasional value approaching K. See Figure 1. Random values from a Zipf distribution can be generated in the manner of [7].

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Algorithm 1: Generate a diverse set of clusterings
Input: X = \{x_1, x_2, ..., x_n\} for x_i \in \mathbb{R}^d, k is the
          number of clusters, m is the number of
         clusterings to be generated
Output: A set of m alternate clusterings of the data
            \{C_1, C_2, ..., C_m\} for which
            C_i: \mathbf{X} \mapsto \{1, 2, ..., k\} is the mapping of each
            point x \in \mathbf{X} to its corresponding cluster
begin
     for i = 1 to m do
          \alpha = rand("uniform", [0 \ \alpha_{max}])
          for j = 1 to d do
              w_i = rand("zipf", \alpha)
          \mathbf{X}_i = \emptyset
         for x \in X do
              x' = x \bigcirc w where \bigcirc is pairwise
              multiplication
              \mathbf{X}_i = \mathbf{X}_i + \{x'\}
          C_i = \text{K-means}(\mathbf{X}_i, k)
     end
```

Algorithm 1 is the procedure that generates different clusterings. First the Zipf shape parameter, α , is drawn uniformly from the interval $[0,\alpha_{max}]$. Here we use $\alpha_{max}=1.5$. (This allows us to sample the space of random weightings, from a uniform distribution ($\alpha=0$) to a severe distribution ($\alpha=1.5$) that gives significant weight to just a few variables.) Then a weight vector $\mathbf{w}\in R^d$ is generated according to the Zipf distribution with that α . Next the features in the original data set are weighted with the weight vector \mathbf{w} . Finally, k-means is used to cluster the feature reweighted data set.

2.1.3 The Problem With Correlated Features

end

Random feature weights may fail to create diverse clusterings in the presence of correlated features: weights given to one feature can be compensated by weights given to other correlated features.

The problem with correlated features can be avoided by applying Principal Component Analysis [9] to the data prior to weighting. Sometimes PCA yields a more interesting set of distance functions by compressing important aspects of the problem into a small set of components. Other times, however, PCA hides important structure. Because of this, we apply random feature weightings both before and after rotating the vector space with PCA. See [6] for a longer version of this paper that includes a thourough description of the use of PCA in meta clustering.

2.1.4 Dealing with Non-vector Data

Feature weighting only works for data in feature-vector format. Data often is available only as pairwise similarities. This problem can be solved using MultiDimensional Scaling (MDS) [11]. MDS transforms pairwise distances to a feature-vector format to which random weights can then be applied. We implement MDS following [11]. In order to avoid local minima and speed the search, we utilize a variable step size and randomize the locations of the points if progress becomes slow or halts.

2.2. Clustering Clusterings at the Meta Level

The methods in the preceding section generate a large, diverse set of candidate clusterings. Usually it is infeasible for a user to examine thousands of clusterings to find a few that are most useful for the application at hand. To avoid overwhelming the user, meta clustering groups similar clusterings together by clustering the clusterings at a meta level. To do this, we need a similarity measure between clusterings.

2.2.1 Measuring the Similarity Between Clusterings

Several measures of clustering similarity have been proposed in the literature [18, 19, 20]. Here we use a measure of clustering similarity related to the Rand index [29]: define I_{ij} as 1 if points i and j are in the same cluster in one clustering, but in different clusters in the other clustering, and I_{ij} is 0 otherwise. The dissimilarity of two clustering models is defined as:

$$\frac{\sum_{i < j} I_{ij}}{N(N-1)/2},$$

where N is the total number of data points. This measure is a metric as well. In the remainder of this paper we refer to it as Cluster Difference.

2.2.2 Agglomerative Clustering at the Meta Level

Once the distances between all pairs of clusterings are found using the Cluster Difference metric, the clusterings are themselves clustered at the meta level. This meta clustering can be performed using any clustering algorithm that works with pairwise similarity data. We use agglomerative clustering at the meta level because it works with similarity data, because it does not require the user to prespecify the number of clusters, and because the resulting hierarchy makes navigating the space of clusterings easier. ([14] presents one of the few studies to examine the tradeoffs between clustering complexity, efficiency, and interpretability.)

An alternate approach for presenting different clusterings to users is to find a low-dimensional projection of the pairwise clustering distances. For example, MDS can be used to find principal components of the meta clustering space from the clustering similarity data. The clusterings can then be presented to the user in a low dimensional plot where similar clusterings are positioned close to each other. One problem we have found with this approach is that often the meta level clustering space is not low in dimension, so any low-dimensional projection of the data has significant distortion. Although easier to visualize, this makes the 2-D projections less useful than the meta clustering tree.

3. Data Sets

Data Set	features	cases	classes	clusters
Australia	17	245	10	10
Bergmark	254	1000	25	25
Covertype	49	1000	7	15
Letters	617	514	7	10
Protein	ud format	639	N/A	20
Phoneme	10	990	15,11	15

Table 1. Description of Data Sets

We evaluate meta clustering on six data sets. For each data set we also classify points using labels external to the clustering. We will call these test classifications the auxiliary labels. The labels are intended as an objective proxy for what users might consider to be good clusterings for their particular application. In practice, users may have only a vague idea of the desired clustering (and thus may not be able to provide the constraints necessary for semisupervised clustering [10, 33]). Or users may have no idea what to expect from the clustering. The auxiliary labels are meant to represent one clustering users might find useful. In no way are they intended to represent an exclusive ground truth for the clustering. If such a classification existed, supervised learning would be more appropriate. Instead, the auxiliary labels are intended to represent a clustering that is good for one application, while acknowledging many applications with other good clusterings exist.

The Australia Coastal data is a subset of the data available from the Biogeoinformatics of Hexacorals environmental database [28]. The data contain measurements from the Australia coastline at every half-degree of longitude and latitude. The features describe environmental properties of each grid cell such as temperature, salinity, rainfall, and soil moisture. Each variable was scaled to have a mean of zero and a mean absolute deviation of one. The auxiliary labels are based on the "Terrestrial Ecoregions of the World" available from [16].

The Bergmark data was collected from 25 focused web crawls, each with different keywords. The variables are

counts in a bags-of-words model for the pages. The auxiliary labels are the 25 web crawls that generated the data.

The Covertype data is from the UCI Machine Learning Repository [27]. It contains cartographic variables sampled at 30×30 meter grid cells in wilderness areas. Data was scaled to a mean of zero and a standard deviation of one. The true forest cover type for each cell is used as the auxiliary labels.

The letters data is a subset of the isolet spoken letter data set from the UCI Machine Learning Repository [27]. We took random utterances of the letters A, B, C, D, F, H, and J in these proportions: A: 109 B: 56 C: 126 D: 59 F: 49 H: 62 J: 53. Each utterance is described by spectral coefficients, contour features, sonarant features, pre-sonarant features, and post-sonarant features. The spoken letters are the auxiliary labels.

The Protein data is the pairwise similarities between 639 proteins. It was created by crystallographers developing techniques to learn relationships between protein sequence and structure. This data is used as a case study in Section 5.

The Phoneme data is from the UCI Machine Learning Repository [27]. It records the 11 phonemes of 15 speakers. This data is used as a case study in Section 6.

See Table 1 for a summary of the data sets.

4. Experimental Results

In this section we present empirical results on four test problems used to develop meta clustering. First, in Section 4.1 we show the effect of Zipf random weighting of feature vectors. In Section 4.2 we compare k-means with weighted feature vectors to the local minima found by k-means on unweighted data. Then in Section 4.4 we show the results of agglomerative hierarchical clustering at the meta level. The results demonstrate the importance of generating a diverse set of clusterings when the clustering objective is not well defined prior to clustering.

We examine two clustering performance metrics. The first is compactness. Compactness is defined as:

$$\frac{\sum_{i=1}^{k} N_i \frac{\sum_{j=1}^{N_i - 1} \sum_{k=j+1}^{N_i} d_{jk}}{N_i (N_i - 1)/2}}{N}, \tag{1}$$

where k is the number of clusters; N_i is the number of points in the i^{th} cluster; d_{jk} is the distance between points j and k, and $N = \sum_{i=1}^k N_i$. Compactness measures the average pairwise distances between points in the same cluster. Regardless of the feature weighting used in clustering, compactness is always measured relative to the original data set. In many traditional clustering algorithms such as kmeans and agglomerative clustering, the optimization criterion is closely related to this measure of compactness.

The second clustering performance metric is accuracy, which is measured relative to the auxiliary labels for each

data set. Again, the auxiliary labels are only a proxy for what a specific user might want in a particular application. They do not represent a single "true" classification of the data, as different users may desire different clusterings for alternate applications. Accuracy is defined as:

$$\frac{\sum_{i=1}^{k} \max(C_i|L_i)}{N},\tag{2}$$

where C_i is the set of points in the i^{th} cluster; L_i is the labels for all points in the i^{th} cluster, and $max(C_i|L_i)$ is the number of points with the plurality label in the i^{th} cluster (if label l appeared in cluster i more often than any other label, then $max(C_i|L_i)$ is the number of points in C_i with the label l).

Determining the number of clusters in a data set is challenging [26]. Indeed, the "correct" number of clusters depends on how the clustering will be used. For simplicity, we make the unrealistic assumption that the desired number of clusters is predefined. It is easy to extend meta clustering to explore the number of clusters as well: in addition to k-means local minima and various variable weighting, clusterings can be generated with different numbers of clusters. Our similarity metric can accommodate clusterings with different numbers of clusters, so the meta-level clustering will still group similar clusterings together, even if they have different numbers of clusters. Using a fixed k helps to demonstrate how effective the methods we present are at generating diverse clusterings.

4.1. Effect of Zipf Weighting

In Figure 2 each point in the plots represents an entire clustering of the data. The scatter plots in each row show clusterings generated by weighting features with Zipf-distributed random weights with different shape parameters α for the four test data sets. We test Zipf distributions with $\alpha = 0.00, 0.25, 0.50, 0.75, 1.00, 1.25, and 1.50.$

Note that for different α values, feature weightings explore different regions of the clustering space. In all data sets that we test, as the α value increases, feature weighting explores a region of lower compactness. We also observe an interesting phenomenon that some of the most accurate clusterings are generated when applying feature weighting with higher α values. In particular, high α in the Covertype data set reveals a cloud of more accurate (and much less compact) clusterings which do not show up at low α values. When $\alpha=0.00$ (1st row) the Zipf distribution yields a uniform distribution. It is clear that a uniform distribution alone is insufficient to explore the clustering space.

The bottom row of Figure 3 shows histograms of the compactness of the clusterings. The most compact clusterings are on the left of the histograms. The arrow in each figure marks the most accurate clusterings. In Australia and Covertype, the most accurate clusterings are not in the top

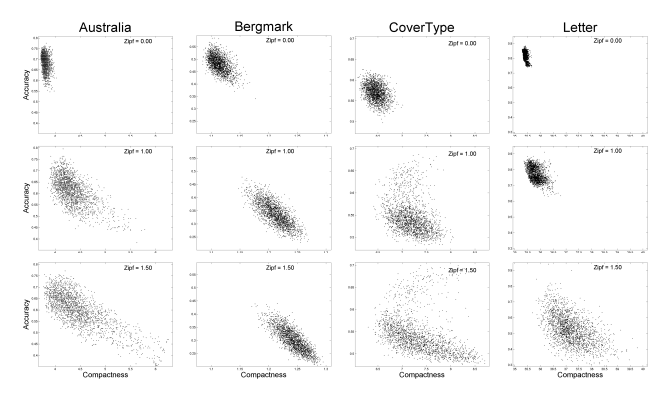


Figure 2. Accuracy vs. compactness for Zipf-weighted sequences for the Australia, Bergmark, Covertype, and Letter data sets. *Each point is a full clustering*.

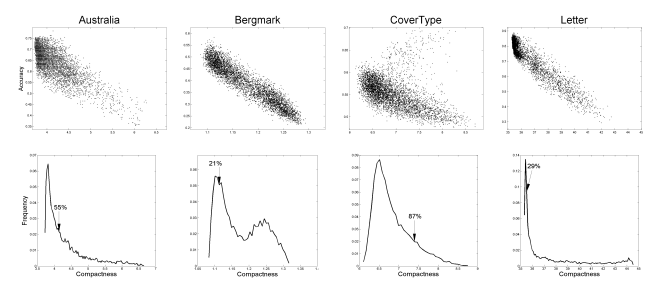


Figure 3. Accuracy vs. compactness for Zipf-weighted clusterings (top) and histograms (bottom) of the compactness of the clusterings. Each point is a full clustering. The arrows indicate the location of the most accurate clustering, and the number above is the percent of clusterings more compact than the most accurate one.

50% of the most compact clusterings. In Bergmark and Letter, the most compact clusterings occur 20-30% down in the distribution. Standard clustering techniques would never find these most accurate clusterings. Looking at the

leftmost points in each plot, it is clear that the most compact clustering is significantly less accurate than the most accurate clustering. Since there exist reasonable clusterings which are more accurate than the most compact clus-

tering, we conclude that the usual method of searching for the most compact clustering can be counterproductive. It is important to explore the space of reasonable clusterings more thoroughly.

4.2. Local Minima vs. Feature Weighting

Figure 4 shows scatter plots of clusterings generated using k-means, comparing weighting features with Zipf distributed random weights versus without using feature weighting for the four test problems. The x-axis is the compactness of the clusterings (as defined in Section 4). The y-axis is the accuracy of the clusterings using the auxiliary labels. The top row shows clusterings generated by random Zipf weighting. The bottom row shows the clusterings generated using iterated k-means where each point is the result of a different random initialization.

For Australia, Bergmark, and Letter, weighting features yields more diverse clusterings – the figures extend further to the lower right, while retaining points in the upper left. For Covertype, not applying feature weighting fails to discover the cloud of more accurate (yet not compact) clusterings. For Australia, different random initialization of kmeans without feature weighting finds more clusterings in the upper left corner (both accurate and compact).

4.3. Other Clustering Methods

For a thorough comparison, we also apply other clustering approaches to the four data sets: hierarchical agglomerative clustering (HAC) [22], EM-based mixture model clustering [23], and two types of spectral clustering [24, 1] (See Figure 4). HAC is implemented using three different linkage criteria: single (min-link), complete (max-link), and centroid (average-link) [22]. The EM-based mixture model clustering estimates Gaussian mixture models using the EM algorithm. The result for the EM-based mixture model clustering is the highest accuracy from multiple runs with different initialization. For both the spectral clustering methods, we also report the highest accuracy from multiple runs. Spectral clustering requires an input similarity matrix S, where $S = exp(-||x_i - x_j||/\sigma^2)$. To obtain a variety of results, we used different values of the shape parameter, $\sigma = \frac{\max\{||x_i - x_j||^2\}}{2^{k/8}}$, for k = 0, ..., 64. The clusterings obtained from other methods fall within the range of clusterings generated from k-means with random feature weighting. Across the four data set the spectral method proposed by [1] has the best performance, though it (and the other alternative clustering methods) are always outperformed by at least one clustering found with the metaclustering technique using kmeans for the base-level clusterings. We are currently investigating random feature weighting with other clustering techniques (in addition to k-means) to create diverse base-level clusterings.

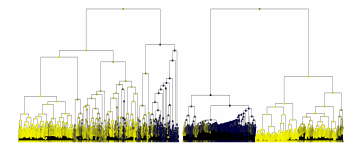


Figure 5. Meta level clustering trees for Australia (left) and Letter (right). Color (yellow to blue) indicates to accuracy (low to high). (The meta clustering dendrograms are much easier to interpret if viewed in color.)

4.4. Meta-Level Agglomerative Clustering

In this section we cluster the clusterings at the meta level using agglomerative clustering with the "flexible strategy" criterion described in [22] and show that clusterings near each other at the meta level are similar, and clusterings far from each other at the meta level are qualitatively different.

Figure 5 shows the meta level clustering trees for two data sets, Australia and Letter. Each node in the clustering tree is colored by accuracy. Yellow indicates low accuracy. Blue indicates high accuracy. Note that clusterings of similar accuracy tend to be grouped together at the meta level even though meta clustering has no information about accuracy. The results suggest that groups of clusterings at the meta level reflect aspects of the base-level clusterings that might be important to users.

Figure 6 shows four representative clusterings of the Australia data set selected from the meta clustering tree. The clusterings on the left were selected as two very similar (but not identical) clusterings from one region of the tree, while the two on the right were selected to be distinct from each other and from the two on the left. Each represents a different, reasonable way of clustering the data set. Once again we see that clusterings close to (far from) each other at the meta level are similar to (different from) each other.

Figure 7 shows compactness (Equation 1) vs. the number of clusters for hierarchical agglomerative clustering at the meta level for the four data sets. These plots can be used to detect structure at the meta level; jumps in the curves indicate merges between groups of dissimilar clusterings. The Australia, Covertype, and Letter data sets exhibit significant structure at the meta level. However, in the Bergmark data set, little structure at the meta level is observed. (Meta clustering can be useful even in the absence of significant structure at the meta level since clustering at the meta level will still group similar clusterings together, making exploring the space of clusterings easier.)

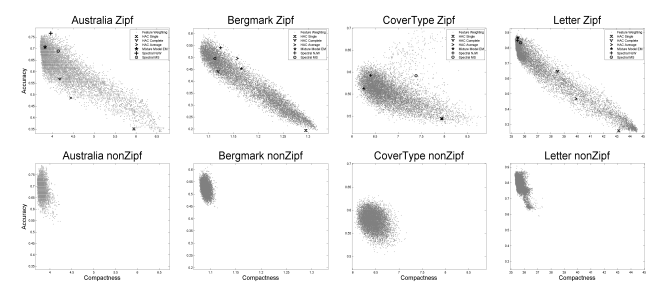


Figure 4. Accuracy vs. compactness for Zipf-weighted clusterings (top row) and iterated k-means (bottom row) of the four data sets: Australia, Bergmark, Cover Type, and Letters. Each point is a full clustering of the data. The top row also includes the accuracy and compactness of other popular clustering methods marked with special symbols.

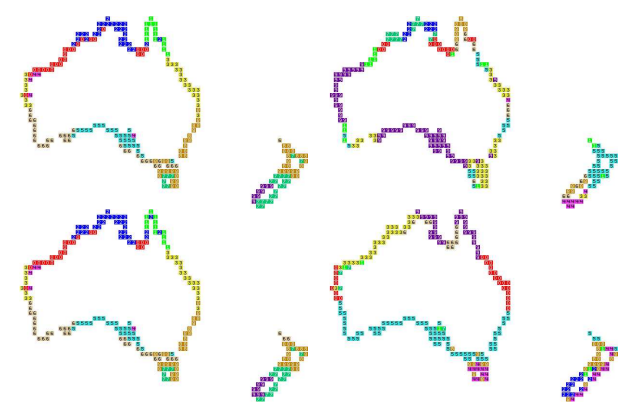


Figure 6. Alternate clusterings for the Australia data set. Each point is colored and numbered by cluster membership. The two clusterings on the left are similar, but not identical, while the two on the right are distinct from each other and from the two on the right. All four are reasonable ways of clustering Australia. (The figures include small numbers that are visible in black and white.)

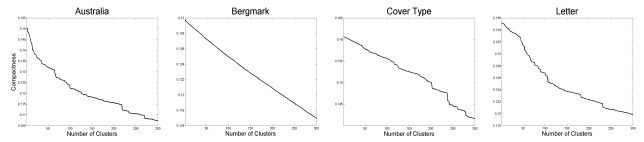


Figure 7. The compactness plot of the hierarchical agglomerative clustering at the meta level.

5. Case Study: Protein Clustering

In this section we apply meta clustering to a real protein clustering problem. The protein data consists of pairwise distances between 639 proteins. The distance between each pair of proteins is computed by aligning 3-D structures of each protein and computing the mean distance in angstroms between corresponding atoms in the two structures.

This data was created by crystallographers developing automated techniques to learn relationships between protein sequence and structure. For their work they need to find groups of proteins containing as many proteins with similar structure as possible. To achieve this goal, and also to better understand the data, the developers employed a number of techniques including clustering, MDS, univariate, and multivariate statistical analysis. In the course of their work they generated and tested a large variety of hand-tuned clustering criteria before finding satisfactory clusterings. After a month of effort, the two largest homogeneous groups of proteins (less than 1.25 angstrom mean distance between aligned atoms) discovered contained 28 and 30 proteins.

We apply meta clustering to the same data to try to automatically find sets of proteins similar to those the experts found manually. For this data we only have pairwise distances between proteins, not a feature-space, so we cannot apply random Zipf weighting directly to the original attributes. Instead, we use the MDS method described earlier in Section 2.1.4 to convert the pairwise distance matrix to a vector space, and then apply random Zipf weighting.

We apply the meta clustering method described in Section 2.1.2 using random Zipf weighting with Zipf shapes selected uniformly on the range (0.0,1.5). For each weighting we run iterated k-means to find clusterings (local minima). We repeat this process 5000 times, yielding 5000 different clusterings of the protein data. The left plot in Figure 8 shows the number of points in the largest cluster satisfying the 1.25 angstrom constraint, plotted as a function of clustering compactness. Note that although the majority of clusterings found by meta clustering do not have large clusters with 30 or more points, meta clustering has found a number of clusterings that contain clusters with more than 30 points. The largest homogeneous cluster found by meta clustering

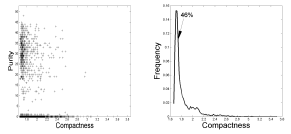


Figure 8. Meta Clustering of the Protein Data

contains 48 proteins, 60% more than the experts were able to find using manually guided methods. In a few hours meta clustering finds better clusterings than could be found manually with a month of work. The compactness histogram in the right of the Figure 8 shows that the "optimal" clustering had mediocre overall compactness, falling near the middle of the distribution of clustering compactnesses.

An examination of compactness (see Section 4.4) as a function of the number of clusters for agglomerative clustering at the meta level shows large jumps, suggesting structure at the meta level, i.e., meta clustering has found qualitatively different ways of clustering the protein data that cluster together at the meta level. If users needed clusterings that satisfied different criteria, it is likely that groups of alternate clusterings have already been found by meta clustering that would perform well according to these other criteria.

The clustering goal used in this case study (clusterings smaller than 1.25 angstroms containing many points) was not known in advance when crystallographers began working with this data. This criterion emerged only after examining the results of many clusterings. Optimizing directly to this criterion is not straightforward. Meta clustering automatically finds a diverse set of clusterings, a few of which have the desired property, without knowing the criterion in advance and without optimizing directly to that criterion.

6. Case Study: Phoneme Clustering

In this section we apply meta clustering to a phoneme clustering problem. The data set contains 15 different speakers saying 11 different phonemes 6 times each (for a total of 990 data points). For this data set, we consider users interested in identifying either speakers or phonemes and evaluate the clusterings based on both of these criteria.

Meta clustering successfully finds clusterings that are accurate for each criterion. Figure 9 shows the scatter plots of clusterings (top two rows) and the meta level clustering dendrograms (bottom row) colored with respect to the two accuracy measures (identifying speakers and recognizing phonemes). In the scatter plots of compactness and accuracy (top row), there is a small cloud of clusterings with high accuracy and medium compactness. If the task were to identify speakers, the most accurate clustering occurs at 38% in the compactness distribution, i.e. the most accurate clusterings. For the task of identifying phonemes, the most accurate clustering does not even occur in the top half of the most compact clusterings and falls at 53% in the compactness distribution.

In the scatter plot of the two accuracy measures (second row), the task of identifying speakers and the task of identifying phonemes are essentially uncorrelated. The most accurate clustering for identifying speakers is generated from applying Zipf weighting with the shape parameter set at 0.25. The most accurate clustering of identifying phonemes is generated from applying Zipf weighting with the shape parameter set at 1.25. This confirms the need to sample a variety of Zipf weighting parameters.

For comparison, the scatter plots in Figure 9 show the consensus clustering found using the cluster aggregation method proposed in [25] (marked with a "+" in the figures). This is the clustering that represents the consensus of *all* found clusterings. As expected, the consensus clustering is very compact (because less compact clusterings often disagree with each other, but the most compact clusterings often agree thus forming a strong consensus). Note, however, that the consensus clustering is not as compact as the most compact clusterings found by meta clustering, and also not very accurate on either the speaker identification or phoneme recognition tasks.

Again we use agglomerative clustering at the meta level to group similar clusterings. Figure 9 shows two copies of the same meta level clustering dendrogram (bottom row) colored by accuracy on the speaker identification and phoneme recognition tasks. Asterisks under the dendrograms indicate groups of clusterings that have significantly different accuracy for the two different tasks. Users may examine different clusterings by clicking on a clustering in the dendrogram, allowing users to zero-in on regions that appear promising.

If a user selects a cluster of clusterings in the meta level dendrogram (as opposed to a single base-level clustering), they can examine either the most central clustering in this branch of the dendrogram, or can examine a consensus clus-

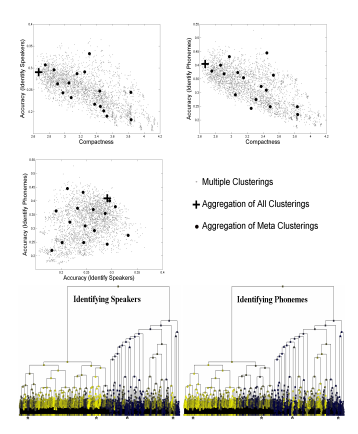


Figure 9. Accuracy vs. compactness scatter plots for the two accuracy measures (identifying speakers vs. recognizing phonemes) (1st row), scatter plot of the two accuracy measures (2nd row), meta level clustering dendrograms shaded by accuracy in the two measures. (The dendrograms are much easier to interpret if viewed in color. See [6])

tering formed from the clusterings in the branch. The large dots in the scatter plots represent consensus clusterings that a user might select.¹

7. Related Works

[2] presents a very different algorithm for finding alternate clusterings of the data. In this approach a probability matrix that defines the likelihood of jumping from one point to another is used to generate a random walk. The transition probability is defined as a function of the Euclidean distance between each pair of points. The random walk allows particles to transition between instances according to the transition probability. Instead of clustering the data directly, distributions of the locations of the particles are clus-

 $^{^1}$ We use the meta level clusterings found for k=16 because examination of compactness vs. number of clusters for the meta-level agglomerative clustering indicated that there were 16 natural meta-level clusters.

tered. Gaps in the eigen values indicate potentially good partitionings. At any random walk step with a local maximum eigen gap, the partition that maximizes this gap is reported. One of the ways in which this approach differs from meta clustering is that it uses a fixed method for measuring the distance between instances (euclidean distance). Also, the method only generates one clustering for each K. (All of the clusterings found with meta clustering in this paper are for a single fixed K.)

A number of ensemble clustering methods improve performance by generating multiple clusterings. We mention only a few here. The cluster ensemble problem is formulated as a graph partitioning problem in [31] where the goal is to encode the clusterings into a graph and then partition it into K parts with the objective of minimizing the sum of the edges connecting those parts.

[12] proposes a different cluster ensemble method where both clusters and instances are modeled as vertices in a bipartite graph. Edges connect instances with clusters with a weight of zero or one depending on whether the instance does or does not belong to the cluster, thus capturing the similarity between instances and the similarity between clusters when producing the final clustering.

Another cluster ensemble method was proposed by [17] where the objective is to find a final clustering that has minimum disagreement between it and all of the clusterings. In this framework, the clustering aggregation problem is mapped to the correlation clustering problem where we have objects and distances between every pair of them and the goal is to produce a partition that minimizes the sum of the distances inside each partition and maximizes the sum of the distances across different partitions.

Instead of partitioning, [25] used agglomerative clustering to produce the final clustering after generating a similarity matrix from many base-level clusterings. Cluster aggregation was formulated as a maximum likelihood estimation problem in [32] where the ensemble is modeled as a new set of features that describe the instances and the final clustering is produced by applying K-means while solving an EM problem. Linear programming was used in [4] to find the relation between the clusters in the different clusterings and the clusters of the final clustering. Simulated annealing and local search was used in [13] to find the final clustering.

The main difference between these ensemble methods and meta clustering is that most ensemble methods combine the clusterings they find into one final clustering because their goal is to find a better, single, very compact clustering. Because the most compact clusterings are not necessarily the most useful clusterings, meta clustering does not combine different clusterings into one clustering. Instead, it groups different clusterings into meta clusters to allow users to select the clustering that is most useful for them.

8. Summary

Searching for the single best clustering may be inappropriate: the clustering that is "best" depends on how the clusters will be used and the data may need to be clustered in different ways for different uses. When clustering is used as a tool to help users understand the data, an appropriate clustering criterion cannot be defined in advance.

The standard approach is for users to try to express in the distance metric a notion of similarity appropriate to the task at hand. This is awkward. Having to define the clustering distance metric, and then refine it when the clusters found are not what was wanted, is akin to having to modify word processing software when the formatting it generates is not what was expected. Few of the many potential users of clustering are adept at defining distance metrics and at understanding the (often subtle) implications a distance metric has on clustering. Even clustering researchers have difficulty modifying distance metrics to achieve better clusterings when the first metric they try does not work adequately.

In this paper we used auxiliary labels not available to clustering to measure clustering accuracy. We use accuracy as a proxy for users who have unspecified goals and intended uses of the clusterings. This allows an objective evaluation of meta clustering. Experiments with four test problems show that meta clustering is able to automatically find superior clusterings. Contrary to common wisdom, in these experiments we find only modest correlation between clustering compactness and clustering accuracy. The most accurate clusterings sometimes are not even in the most compact 50% of the clusterings. This reinforces our belief that searching for a single, optimal clustering is inappropriate when correct clustering criteria cannot be specified in advance. Instead, it is more productive to focus clustering on finding a large number of good, qualitatively different clusterings and allow users (or some form of post processing) to select the clusterings that appear to be best.

Experiments with a phoneme clustering problem showed that clusterings that are good for one criterion can be very suboptimal for another criterion. Different users may need different clusterings. Meta clustering automatically found good (different) clusterings for each criterion. Experiments with a protein clustering problem provide a case study where meta clustering was able to improve clustering quality 60% above the best that could be achieved by human experts working with this data. Meta clustering achieved this improvement fully automatically in less than a day of computation. We believe the results demonstrate that meta clustering can make clustering more useful to non-specialists and will reduce the effort required to find excellent clusterings that are appropriate for the task at hand.

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