Adaptive quality-based clustering of gene expression profiles

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ABSTRACT

Motivation: Microarray experiments generate a considerable amount of data, which analyzed properly help us gain a huge amount of biologically relevant information about the global cellular behaviour. Clustering (grouping genes with similar expression profiles) is one of the first steps in data analysis of high-throughput expression measurements. A number of clustering algorithms have proved useful to make sense of such data. These classical algorithms, though useful, suffer from several drawbacks (e.g. they require the predefinition of arbitrary parameters like the number of clusters; they force every gene into a cluster despite a low correlation with other cluster members). In the following we describe a novel adaptive quality-based clustering algorithm that tackles some of these drawbacks.

Results: We propose a heuristic iterative two-step algorithm: First, we find in the high-dimensional representation of the data a sphere where the 'density' of expression profiles is locally maximal (based on a preliminary estimate of the radius of the cluster—quality-based approach). In a second step, we derive an optimal radius of the cluster (adaptive approach) so that only the significantly co-expressed genes are included in the cluster. This estimation is achieved by fitting a model to the data using an EM-algorithm. By inferring the radius from the data itself, the biologist is freed from finding an optimal value for this radius by trial-and-error. The computational complexity of this method is approximately linear in the number of gene expression profiles in the data set. Finally, our method is successfully validated using existing data sets.

Availability: http://www.esat.kuleuven.ac.be/~thijs/Work/Clustering.html

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Supplementary information: http://www.esat.kuleuven.ac.be/~fdesmet/paper/adaptpaper.html

INTRODUCTION

A variety of techniques (Self-Organizing Maps; Tamayo et al., 1999), hierarchical clustering (Carr et al., 1997; Eisen et al., 1998), Self-Organizing Tree Algorithm (Herrero et al., 2001), K-means (Tavazoie et al., 1999; Tou and Gonzalez, 1979), simulated annealing (Lukashin and Fuchs, 2001), Principal Component Analysis (Quackenbush, 2001), MultiDimensional Scaling (Bittner et al., 2000), Cluster Affinity Search Technique (Ben-Dor et al., 1999) has been implemented and successfully been used to analyze or cluster high-dimensional microarray data (DeRisi et al., 1997; Lander, 1999; Schena et al., 1995).

One of the objectives of these methods is to detect groups of genes that exhibit a highly similar expression profile (here defined as coexpressed). Since gene expression profiles are encoded in real vectors (whose elements are the different measurements of the expression levels of a specific gene), these algorithms intend to group gene expression vectors that are sufficiently close to each other (according to a certain distance or similarity measure). Most clustering algorithms originate from non-biologically related research fields. Therefore, although useful, the original implementations suffer from some drawbacks as has been highlighted by Sherlock (2000). These deficiencies can be summarized as follows. Firstly, algorithms such as K-means and Self-Organizing Maps require the predefinition of the number of clusters (parameter of the algorithm). When clustering gene expression profiles, the number of clusters present in the data is usually unknown. Changing this parameter usually affects the final clustering result considerably. Clustering, using for example K-means, therefore involves extensive parameter fine-tuning to detect the optimal clustering and the choice of the final parameter setting remains somehow arbitrary (e.g. based on visual inspection of the clusters). When using hierarchical clustering, the number of clusters is determined by cutting the tree structure at a certain level. The resulting cluster structure is therefore highly influenced by the choice of this level, which in turn is rather arbitrary. Secondly, the idea of forcing each gene of the data set into a cluster is a significant drawback of these implementations. If genes are, despite a rather low correlation with other cluster members, forced to end up in one of
the clusters, the average profile of this cluster is corrupted and the composition of the cluster becomes less suitable for further analyses (such as motif finding or functional annotation: Roth et al., 1998; van Helden et al., 2000).

Much effort is currently being done to adapt clustering algorithms towards the specific needs of biological problems. In this context the ideas of quality-based clustering (Heyer et al., 1999) and gene shaving (Hastie et al., 2000) were developed (gene shaving also uses a quality measure to select the cluster size). Heyer et al. (1999) proposed an algorithm (QT_Clust) that tries to identify clusters that have a certain quality (representing the minimal degree of coexpression needed—see below for the exact definition used in this paper) and where every cluster contains a maximal number of points. Genes not exhibiting this minimal degree of coexpression with any of the clusters are excluded from further analysis. A problem with the quality-based approach of Heyer et al. (1999), however, is that this quality is a user-defined parameter that is hard to estimate (it is hard to find a good trade-off or optimal value: setting the quality too strictly will exclude a considerable number of coexpressed genes, setting it too loose will include too many genes that are not really coexpressed). Moreover, it should be noted that the optimal value for this quality is, in general, different for each cluster and data set dependent.

In this paper, we describe an adaptive quality-based clustering method starting from the principles described by Heyer et al. (1999; quality-based approach; locating clusters, with a certain quality, in a volume where the density of points is maximal). The algorithm described below is in essence a heuristic, two-step approach that defines the clusters sequentially (the number of clusters is not known in advance, so it is not a parameter of the algorithm). The first step locates a cluster (quality-based approach) and the second step derives the quality of this cluster from the data (adaptive approach). The performance of the algorithm is tested on a real microarray data set and the result is compared with an already published analysis (K-means) using the same data. Finally, we make a theoretical comparison between our algorithm and the algorithm of Heyer et al. (1999).

**METHODS**

**Normalization**

It is common practice to normalize gene expression vectors before cluster analysis. In this paper, we normalize the expression profiles so that their mean is zero and their variance is one before proceeding with the actual cluster algorithm. If \( g_i = (g_i^1, g_i^2, \ldots, g_i^E) \) is a normalized expression vector, this means that

\[
\mu_i = \frac{1}{E} \sum_{j=1}^{E} g_i^j = 0.
\]  

Normalized expression profiles or vectors therefore are located in an \( E \)-dimensional space on the intersection of a hyperplane (Equation 1) and a hypersphere with a radius equal to \( \sqrt{E - 1} \) (Equation 2).

**Quality R of a cluster**

The definition used in this paper for the quality \( R \) of a cluster, is as follows: In a collection of gene expression profiles \( G = \{g_i, i = 1, \ldots, N\} \), a cluster \( K \) with center \( C_K \) and quality \( R_K \) (also called radius of cluster \( K \)), will only contain the profiles satisfying the following property:

\[
\|g_i - C_K\|_2 < R_K. \tag{3}
\]

Equation (3) means that cluster \( K \) only contains gene expression profiles with a minimum degree of coexpression (represented by the quality guarantee \( R_K \)). The norm or distance measure we use in this paper is the 2-norm or Euclidean distance.

**ALGORITHM AND IMPLEMENTATION**

**Global algorithm**

The global cluster algorithm (Adap_Cluster) is, as mentioned previously, a heuristic iterative two-step approach where the basic steps are as described below. In this implementation we use two user-defined parameters (MIN_NR_GENES and S—the values between brackets are default values), several internal tuning parameters that have a fixed value (the user is not allowed to change these values) and the data set itself (\( G \)).

\[
\begin{align*}
\text{Adap_Cluster} & \quad (G = \{g_i, i = 1, \ldots, N\}, \text{MIN}_\text{NR}_\text{GENES} < 2, S < 0.95) \\
\text{ACCUR}_\text{RAD} & = 0.1 \quad */ \text{Set internal tuning parameter} */ \\
\text{Initialize} R_K_\text{PRELIM} & \quad /* \text{Radius estimate initialization} */ \\
\text{WHILE} \quad \text{Stop criterion NOT TRUE} \\
\text{END WHILE} \\
\end{align*}
\]

During each iteration, this algorithm first finds a cluster center (\( C_K \)) using a preliminary estimate (\( R_K_\text{PRELIM} \)) and quality

\[
\sigma_i = \sqrt{\frac{1}{E-1} \sum_{j=1}^{E} (g_i^j - \mu_i)^2} = 1. \tag{2}
\]
of the radius or quality of the cluster (Step 1). When the cluster center has been located, the algorithm determines a new estimate for the radius \((R_K)\) of the cluster (Step 2). Now there are two possibilities:

1. If this new estimate is approximately equal to the preliminary estimate (e.g. within 10\%—ACCUR_RAD), the set of genes defined by the cluster center and the new estimate of the radius is removed from the data set \(G\). Furthermore, if the number of genes in this set is equal or larger than a predefined value (MIN_NR_GENES—user-defined; default 2), this set is a valid cluster. The preliminary estimate of the radius to be used in Step 1 of the next iteration (for the next cluster) is updated with the new estimate of the radius calculated in Step 2 of the current iteration (in most cases, the best preliminary estimate for the radius of the next cluster seems to be the radius of the previous cluster).

2. If the new estimate of the radius is substantially different from the preliminary estimate, the preliminary estimate \(R_K\) PRELIM is also updated with the new estimate \(R_K\) and a new iteration is started. This is repeated until the relative difference between \(R_K\) and \(R_K\) PRELIM falls under ACCUR_RAD.

The iterations are terminated when the stop criterion is satisfied (see below).

The algorithm was implemented in MATLAB and is publicly available for data analysis. Note that this implementation can deal with missing values often occurring in expression data (also see Troyanskaya et al., 2001). We used the methodology suggested by Kaufman and Rousseuw (1990) to handle missing values. A detailed mathematical description of the missing values management can be found on our supplementary web site. Below we will discuss the initialization of the preliminary estimate of the radius before the first iteration, the procedures used in Step 1 and 2, the stop criterion (WHILE loop) and the computational and memory complexity of the overall algorithm.

Radius estimate initialization

In the global cluster algorithm, the preliminary estimate of the radius \((R_K\) PRELIM) has to be initialized before the first iteration (radius estimate for the first cluster—line 3 of Adap_Cluster). We use half of the radius of the hypersphere defined by normalization of the expression profiles (see above). This is given by:

\[
R_K\text{ PRELIM} = \sqrt{E - 1 \over 2}
\]  

(4)

where \(E\) is the dimension of the gene expression vectors (number of expression vector components).

Localization of a cluster center—quality-based clustering (Step 1)

Given a collection \(G\) of gene expression profiles, the objective of Step 1 is to find a cluster center in an area of the data set where the ‘density’ (or number) of expression profiles, within a sphere with radius or quality equal to \(R_K\) PRELIM (preliminary estimate of the radius), is locally maximal. The method described here is based on the principles used by Heyer et al. (1999) but is significantly faster (also see Discussion—Tables 2a, b). The disadvantage with the approach of Heyer et al. (1999) is that the quality or radius of the clusters is a parameter that is not very intuitive (it is often hard to find a ‘good’ value for this parameter; often a trial-and-error approach is used with manual validation of the clusters). Furthermore, all the clusters are forced to have the same radius.

The basic steps of the algorithm used for the first step are described below:

\[
C_K = \text{locate_cluster_center}(G, R_K\text{ PRELIM})
\]

\[
\text{MAXITER}=50
\]

/* Set Internal tuning parameter — maximum number of iterations */

\[
\text{DIV}=1/30
\]

/* Set internal tuning parameter — fraction needed to determine DELTRAD */

\[
C_K = \text{mean}(G)
\]

/* Cluster center initialization */

\[
R_K = \text{max}(|\|g_i - C_K\| | g_i \in G) / \|R_K\| \text{ PRELIM}"
\]

/* Start with maximal radius */

\[
\text{DELTARAD} = (R_K - R_K\text{ PRELIM}) \times \text{DIV}
\]

/* Determine step for decreasing radius */

\[
R_K = \text{RAD} - \text{DELTRAD}
\]

/* Decrease radius */

\[
\text{GENES_IN_SPHERE} = \{g_i \in G | |g_i - C_K| < R_K\}
\]

/* Determine profiles within sphere */

\[
\text{ME} = \text{mean} (\text{GENES_IN_SPHERE})
\]

/* Recalculate mean */

\[
\text{ITER} = 1
\]

WHILE (\(\text{ME} \neq C_K\) AND \(\text{ITER} < \text{MAXITER}\) OR \(\text{RAD} > R_K\) PRELIM)

/* Iterate until convergence or maximal number of iterations has been reached */

\[
\text{ITER} = \text{ITER} + 1
\]

\[
C_K = \text{ME}
\]

/* Move cluster center to cluster mean */

IF \(\text{RAD} > R_K\) PRELIM

\[
\text{RAD} = \text{RAD} - \text{DELTRAD}
\]

/* Decrease radius if desired quality has not yet been reached */

END IF

\[
\text{GENES_IN_SPHERE} = \{g_i \in G | |g_i - C_K| < R_K\}
\]

/* Determine profiles within sphere */

\[
\text{ME} = \text{mean}(\text{GENES_IN_SPHERE})
\]

/* Re-calculate mean */

END WHILE

IF \(\text{ME} \neq C_K\)

\[
C_K = \text{empty}
\]

/* Undefined cluster center if no convergence */

END IF

After initialization of the cluster center (with the mean profile of all the expression profiles in the data set \(G\)), all the expression profiles within a sphere with radius RAD are selected. Iteratively, the mean profile of these expression profiles is calculated and subsequently the cluster center is moved to this mean profile. This approach moves the cluster in the direction where the ‘density’ of profiles is higher (conceptually visualized in Fig. 1).

The radius RAD of the sphere is initialized so that all profiles in the data set are located within this sphere.
Table 1a. Biological validation of the cluster result in Figure 3 (see supplementary web site) and comparison with the result of Tavazoie et al. (1999)

<table>
<thead>
<tr>
<th>Cluster number</th>
<th>Number of ORFs</th>
<th>MIPS functional category</th>
<th>ORFs within functional category</th>
<th>P-value ($-\log_{10}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adap_Cluster</td>
<td>Adap_Cluster</td>
<td>K-means</td>
<td>Adap_Cluster</td>
<td>K-means</td>
</tr>
<tr>
<td>(Tavazoie et al.)</td>
<td>(Tavazoie et al.)</td>
<td>(Tavazoie et al.)</td>
<td>(Tavazoie et al.)</td>
<td>(Tavazoie et al.)</td>
</tr>
<tr>
<td>1</td>
<td>1</td>
<td>302</td>
<td>164</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Adap_Cluster</td>
<td>K-means</td>
<td>Adap_Cluster</td>
<td>K-means</td>
</tr>
<tr>
<td>(Tavazoie et al.)</td>
<td>(Tavazoie et al.)</td>
<td>(Tavazoie et al.)</td>
<td>(Tavazoie et al.)</td>
<td>(Tavazoie et al.)</td>
</tr>
<tr>
<td></td>
<td>Adap_Cluster</td>
<td>K-means</td>
<td>Adap_Cluster</td>
<td>K-means</td>
</tr>
<tr>
<td>(Tavazoie et al.)</td>
<td>(Tavazoie et al.)</td>
<td>(Tavazoie et al.)</td>
<td>(Tavazoie et al.)</td>
<td>(Tavazoie et al.)</td>
</tr>
<tr>
<td>Ribosomal proteins</td>
<td>101</td>
<td>64</td>
<td>80</td>
<td>54</td>
</tr>
<tr>
<td>Organization of cytoplasm</td>
<td>146</td>
<td>79</td>
<td>77</td>
<td>39</td>
</tr>
<tr>
<td>Protein synthesis</td>
<td>119</td>
<td>NR</td>
<td>74</td>
<td>NR</td>
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<tr>
<td>Cellular organization</td>
<td>211</td>
<td>NR</td>
<td>34</td>
<td>NR</td>
</tr>
<tr>
<td>Translation</td>
<td>17</td>
<td>NR</td>
<td>9</td>
<td>NR</td>
</tr>
<tr>
<td>Organization of chromosome structure</td>
<td>4</td>
<td>7</td>
<td>1</td>
<td>4</td>
</tr>
<tr>
<td>Mitochondrial organization</td>
<td>62</td>
<td>32</td>
<td>18</td>
<td>10</td>
</tr>
<tr>
<td>Energy</td>
<td>35</td>
<td>NR</td>
<td>8</td>
<td>NR</td>
</tr>
<tr>
<td>Protein synthesis</td>
<td>24</td>
<td>NR</td>
<td>4</td>
<td>NR</td>
</tr>
<tr>
<td>Ribosomal proteins</td>
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<td>NR</td>
<td>4</td>
<td>NR</td>
</tr>
<tr>
<td>Protein destination</td>
<td>49</td>
<td>NR</td>
<td>4</td>
<td>NR</td>
</tr>
<tr>
<td>DNA synthesis and replication</td>
<td>20</td>
<td>23</td>
<td>18</td>
<td>16</td>
</tr>
<tr>
<td>Cell growth, cell division and DNA synthesis</td>
<td>48</td>
<td>NR</td>
<td>17</td>
<td>NR</td>
</tr>
<tr>
<td>Recombination and DNA repair</td>
<td>12</td>
<td>11</td>
<td>8</td>
<td>5</td>
</tr>
<tr>
<td>Nuclear organization</td>
<td>32</td>
<td>40</td>
<td>8</td>
<td>4</td>
</tr>
<tr>
<td>Cell-cycle control and mitosis</td>
<td>20</td>
<td>30</td>
<td>7</td>
<td>8</td>
</tr>
<tr>
<td>Mitochondrial organization</td>
<td>15</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Peroxisomal organization</td>
<td>4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Energy</td>
<td>9</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TRNA-synthetases</td>
<td>14</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Organization of cytoplasm</td>
<td>14</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cellular transport and transport mechanisms</td>
<td>6</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Transcription</td>
<td>9</td>
<td>21</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>Organization of centrosome</td>
<td>3</td>
<td>6</td>
<td>4</td>
<td>6</td>
</tr>
<tr>
<td>Nuclear biogenesis</td>
<td>1</td>
<td>3</td>
<td>2</td>
<td>5</td>
</tr>
<tr>
<td>Organization of cytoskeleton</td>
<td>2</td>
<td>7</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>TRNA transcription</td>
<td>3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Organization of cytoplasm</td>
<td>7</td>
<td>30</td>
<td>6</td>
<td>9</td>
</tr>
<tr>
<td>Ribosomal proteins</td>
<td>4</td>
<td>16</td>
<td>4</td>
<td>7</td>
</tr>
<tr>
<td>Protein synthesis</td>
<td>4</td>
<td>20</td>
<td>3</td>
<td>7</td>
</tr>
<tr>
<td>Cellular organization</td>
<td>7</td>
<td>55</td>
<td>2</td>
<td>5</td>
</tr>
<tr>
<td>Organization of endoplasmatic reticulum</td>
<td>4</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Every iteration, this radius is decreased with a constant value (DELTARAD, a fraction (DIV) of the difference between the initial value of RAD and $R_K_{PRELIM}$) until the radius has reached the desired value ($R_K_{PRELIM}$) and then remains constant. In the first iterations (when RAD is still ‘large’) this technique will move the cluster center to regions of the data where the ‘global’ density is higher (these regions often contain the largest cluster(s)). After some iterations (when RAD is equal or close to $R_K_{PRELIM}$) the cluster center will move towards an actual cluster where the density is ‘locally’ higher.
Adaptive quality-based clustering

Table 1a. — Continued.

<table>
<thead>
<tr>
<th>Cluster number</th>
<th>Number of ORFs</th>
<th>MIPS functional category</th>
<th>ORFs within functional category</th>
<th>P-value ($-\log_{10}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>42</td>
<td>12</td>
<td>Cellular transport and transport mechanisms</td>
<td>6</td>
<td>4</td>
</tr>
<tr>
<td>5</td>
<td>152</td>
<td>Cell rescue, defense, cell death</td>
<td>22</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Carbohydrate metabolism</td>
<td>24</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Stress response</td>
<td>12</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Energy</td>
<td>16</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Metabolism of energy reserves</td>
<td>6</td>
<td>4</td>
</tr>
<tr>
<td>7</td>
<td>101</td>
<td>Cell-cycle control and mitosis</td>
<td>17</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Budding, cell polarity, filament formation</td>
<td>10</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>DNA synthesis and replication</td>
<td>7</td>
<td>4</td>
</tr>
<tr>
<td>8</td>
<td>148</td>
<td>TCA pathway</td>
<td>5</td>
<td>4</td>
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<tr>
<td></td>
<td></td>
<td>Carbohydrate metabolism</td>
<td>22</td>
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<td>21</td>
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<td>Protein synthesis</td>
<td>14</td>
<td>5</td>
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<tr>
<td></td>
<td></td>
<td>Organization of cytoplasm</td>
<td>18</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Ribosomal proteins</td>
<td>10</td>
<td>4</td>
</tr>
<tr>
<td>30</td>
<td>60</td>
<td>Nitrogen and sulphur metabolism</td>
<td>9</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Amino acid metabolism</td>
<td>12</td>
<td>7</td>
</tr>
</tbody>
</table>

The genes in each cluster have been mapped to the functional categories in the MIPS database and ($-\log_{10}$ transformed) P-values (representing the degree of enrichment—also see text) have been calculated for each functional category in each cluster. Only significantly enriched functional categories are shown ($\log_{10}$ transformed P-values $\geq 4$) and clusters without enrichment are not listed. As a comparison and in parallel (functionally matching clusters are shown in the same row), the results obtained by Tavazoie et al. (1999) (K-means) are also included. NR = Not Reported.

Convergence is reached if the cluster center remains stationary after RAD has reached $R_K_{\text{PRELIM}}$. If this does not happen within a certain (MAXITER) number of iterations, $C_K$ is emptied and the algorithm stops.

Note that, the number of distance calculations performed during each iteration of locate_cluster_center is equal to the number ($= N$) of all expression profiles in $G$ (only the distances from the expression profiles to the current cluster center have to be calculated). Note also that the computational complexity of the calculation of one distance is $O(E)$ ($E$ is the dimensionality of the expression vectors). Because the number of iterations is limited (MAXITER), the computational complexity for the localization of one cluster center is thus $O(N \times E)$.

Re-estimation or adaptation of the cluster quality (Step 2)

In the previous paragraph we located a cluster center $C_K$ in a collection $G$ of gene expression profiles, using a preliminary estimate $R_K_{\text{PRELIM}}$ of the radius of the cluster. The objective of the method described in this paragraph is, given the cluster center that remains fixed, to re-calculate the radius $R_K$ of the current cluster as to assess that genes belonging to this cluster are significantly coexpressed.

To substantiate the method described here, we introduce a randomized version of the original data set where the components of each expression vector are randomly and independently permuted (Herrero et al., 2001). This randomized version of the data will only be used for conceptual reasons, it will not be used during the actual calculations. This process of randomization destroys the correlation between the expression vectors that was introduced through non-accidental mechanisms (e.g. experimental setup). Any correlation still existing after this procedure can be attributed to chance.

First, we calculate the Euclidean distance $r$ from every expression vector in the data set to the cluster center $C_K$. Imagine doing the same for every vector present in the randomized data. The distribution of these distances in the original data consists of two parts (this approach has some similarities with the work of Sharan and Shamir (2000))—see Figure 2:
Table 1b. Biological validation of the cluster analysis of the Cho et al. (1998) data with Adap_Cluster (MIN_NR GENES = 10, S = 0.95) using the standard deviation (σ) as the metric of variance for filtering. The algorithm retrieved 38 clusters. We looked for enrichment (represented by the P-values) of top-level functional categories (from the MIPS database) in individual clusters. Notice the periodic behaviour of the clusters enriched with cell-cycle specific genes (cluster 3, 6 and 9).

<table>
<thead>
<tr>
<th>Cluster number</th>
<th>Graphical representation of cluster</th>
<th>Number of ORFs</th>
<th>MIPS functional category (top-level)</th>
<th>ORFs within functional category</th>
<th>P-value (−log_{10})</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td><img src="image1" alt="Graph" /></td>
<td>426</td>
<td>energy transport facilitation</td>
<td>47</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>40</td>
<td>5</td>
</tr>
<tr>
<td>3</td>
<td><img src="image2" alt="Graph" /></td>
<td>196</td>
<td>cell growth, cell division and DNA synthesis</td>
<td>48</td>
<td>5</td>
</tr>
<tr>
<td>4</td>
<td><img src="image3" alt="Graph" /></td>
<td>149</td>
<td>protein synthesis and cellular organization</td>
<td>71</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>107</td>
<td>19</td>
</tr>
<tr>
<td>5</td>
<td><img src="image4" alt="Graph" /></td>
<td>159</td>
<td>cell rescue, defense, cell death and ageing</td>
<td>20</td>
<td>4</td>
</tr>
<tr>
<td>6</td>
<td><img src="image5" alt="Graph" /></td>
<td>171</td>
<td>cell growth, cell division and DNA synthesis</td>
<td>76</td>
<td>24</td>
</tr>
<tr>
<td>9</td>
<td><img src="image6" alt="Graph" /></td>
<td>78</td>
<td>cell growth, cell division and DNA synthesis</td>
<td>23</td>
<td>4</td>
</tr>
<tr>
<td>37</td>
<td><img src="image7" alt="Graph" /></td>
<td>11</td>
<td>metabolism</td>
<td>9</td>
<td>6</td>
</tr>
</tbody>
</table>

1. Background: these are the expression profiles with a distance to the cluster center that is also significantly present in the distance distribution of the randomized data set. Genes belonging to the background of the current cluster center either do not belong to any cluster (noise; are not significantly coexpressed with other genes) or belong to another cluster. Genes belonging to other clusters (if not too dominant) will not significantly show up in the distance distribution for the current cluster center (they ‘drown’ in the noise or background).

2. Cluster: these are the expression profiles with a distance to the cluster center that is not significantly present in the distance distribution of the randomized data set (left-sided tail in the distribution of the original data set). Genes belonging to the cluster are significantly coexpressed.
To calculate the true radius of the cluster we need to construct a model (probability density estimation) describing the total distribution of the distance \( r \) in the original data. We propose the following model structure:

\[
p(r) = P_C \cdot p(r|C) + P_B \cdot p(r|B)
\]

(5)

where

\[
p(r|C) = \frac{S_D}{(2\pi \sigma^2)^{D/2}} r^{D-1} \exp \left( -\frac{r^2}{2\sigma^2} \right)
\]

(6)\nonumber

\[
p(r|B) = \frac{S_D}{S_{D+1} (D+1)^{D/2}} r^{D-1}
\]

(7)\nonumber

\[
P_C + P_B = 1
\]

(8)\nonumber

and

\[
D = E - 2
\]

(9)\nonumber

\[
S_D = \frac{2\pi^{D/2}}{\Gamma(D/2)}
\]

(10)\nonumber

\[
\Gamma(x) = \int_0^\infty u^{x-1} e^{-u} du.
\]

(11)\nonumber

\( E \) is the dimensionality of the gene expression vectors, \( S_D \) is the surface area of a unit sphere in \( D \) dimensions and \( \Gamma(x) \) is the gamma function.

Note that the model structure assumes that the distance measure used for \( r \) is the Euclidean distance. This means that our method cannot be directly extrapolated to other distance measures.

The model for the total distribution described in Equation (5) is the sum of two terms (also see Figure 2). One term represents the distribution of the cluster (see Equation 6), the other term represents the distribution of the background (see Equation 7), each multiplied by the associated a priori probability (\( P_C \) and \( P_B \)). Note that Equations (6) and (7) are only valid for normalized gene expression vectors. Note also that this model is an approximation and only reliable in the neighbourhood of the cluster. A detailed mathematical discussion of Equations (6) and (7) and of the assumptions used to construct them, can be found on our supplementary website. Notice that two parameters (\( \sigma \) and \( P_C(\text{or } P_B) \)) still have to be determined by fitting the model to the distance distribution of the original data (the randomized data is not used for the actual calculations). This is done by an EM-algorithm (Bishop, 1995). We use the preliminary estimate of the radius \( R_{K, \text{PRELIM}} \) (see Equation 5) to initialize the two parameters to be determined by the EM-algorithm. Because the model only has to fit the distribution of \( r \) (distance to the cluster center—one dimension), the computational complexity of the EM-algorithm is low as compared to the computational complexity of the cluster center localization in Step 1 and therefore can be neglected if \( E \) is sufficiently large. The accuracy of the fit (which represents the validity of the assumptions we made to construct our model) for the clusters found in the Cho et al. (1998) data (see Figure 3 on our supplementary website and the Section Results) can be inspected in Figure 2 for the first four clusters of Figure 3 and on our supplementary website for all the clusters of Figure 3.

After the estimation of \( \sigma \) and \( P_C \), we determine the radius \( R_K \) of the current cluster so that points that are assigned to the cluster have a probability \( S \) or more (significance level—user-defined; default setting: \( S = 95\% \)) to belong to the cluster:

\[
P(C|R_K) = \frac{P_C \cdot p(R_K|C)}{P_C \cdot p(R_K|C) + P_B \cdot p(R_K|B)} = S \quad (12)
\]

To summarize, the complete input–output relation of the method explained in this paragraph is given by

\[ R_K = \text{recalculate radius}(G, C_K, R_{K, \text{PRELIM}}, S). \]

\( R_K \) will be empty if \( C_K \) is empty (cluster center localization did not converge) or if the EM-algorithm to determine the model parameters did not converge.

**Stop criterion**

The iteration (WHILE loop) in the global algorithm ends when the stop criterion is satisfied. This is the case when one of the three following conditions holds true:

**Fig. 1.** Conceptual visualization of cluster center \( X_{C_K} \) relocation to the cluster mean \( X_{M_E} \) in two dimensions (one iteration—cluster radius constant—data not normalized). The number of profiles (black dots) within the sphere after relocation is substantially higher than the number of profiles before relocation.
Fig. 2. Distribution of $r$ (distance from expression vectors to a certain cluster center) for the first 4 clusters found in the data set from Cho et al. (1998), (see Figure 3 on our supplementary web site). In each box, the histogram on the left represents the distribution of $r$ in the actual data and the histogram on the right represents the distribution of $r$ in the randomized data (note that the cluster center for the randomized distribution is identical to the cluster center for the actual distribution—the randomization is not applied to the cluster center itself). For each cluster, the model (see Equations (5)-(11)) fitted by the EM-algorithm is superposed on the distribution of the actual data (after multiplication with an appropriate factor to fit the scale (this accounts for the bin size and the number of expression profiles in the data set)). The model for the background (see Equation (7)) is also superposed on the distribution of the randomized data.

Table 2a. Comparison between Adap_Cluster and QT_Clust

<table>
<thead>
<tr>
<th>Adap_Cluster</th>
<th>QT_Clust</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>User-defined parameters</strong></td>
<td>1. Data set $G$</td>
</tr>
<tr>
<td>2. Significance level $S$</td>
<td>2. Quality (radius $R$ or diameter $d$)</td>
</tr>
<tr>
<td>3. Minimum number of genes</td>
<td>3. Minimum number of genes</td>
</tr>
<tr>
<td><strong>MIN_NR_GENES</strong></td>
<td>(termination criterion)</td>
</tr>
<tr>
<td><strong>Computational complexity</strong></td>
<td>$\sim O(N \times E \times VC)$</td>
</tr>
<tr>
<td><strong>Cluster radius $R$</strong></td>
<td>Automatically calculated for each cluster separately—not constant</td>
</tr>
<tr>
<td><strong>Quality measure</strong></td>
<td>Constant and user-defined</td>
</tr>
<tr>
<td>Significance level $S$: statistical parameter that can be chosen independently of data set (default value (0.95) almost always gives meaningful results)</td>
<td>Arbitrary parameter that has to be set by the user in function of a specific data set, after visual inspection of clusters formed at different quality-levels (optimal value is not straightforward)—no meaningful default value</td>
</tr>
<tr>
<td><strong>Number of clusters</strong></td>
<td>Not predefined</td>
</tr>
<tr>
<td><strong>Inclusion of all genes in clusters</strong></td>
<td>No</td>
</tr>
<tr>
<td><strong>Result</strong></td>
<td>Deterministic</td>
</tr>
<tr>
<td></td>
<td>No</td>
</tr>
</tbody>
</table>
### Table 2b. Comparison between Adap_Cluster, hierarchical clustering, Self-Organizing Maps and K-means

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Format of result</strong></td>
<td>Set of clusters</td>
<td>Tree structure difficult to interpret for large data sets</td>
<td>Set of predefined number of clusters</td>
<td>Set of predefined number of clusters</td>
</tr>
<tr>
<td><strong>Principal user-defined parameter</strong></td>
<td>Significance level S</td>
<td>–</td>
<td>Number of clusters / Node topology</td>
<td>Number of clusters</td>
</tr>
<tr>
<td><strong>Additional requirements from the user</strong></td>
<td>Limited (fine-tuning of S is rarely necessary)</td>
<td>Definition of (an arbitrary) level where the tree structure has to be cut</td>
<td>Extensive parameter fine-tuning (comparison of several runs with different parameter settings) is almost always necessary</td>
<td>Extensive parameter fine-tuning (comparison of several runs with different parameter settings) is almost always necessary</td>
</tr>
<tr>
<td><strong>Statistical definition of clusters</strong></td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td><strong>Inclusion of all genes in clusters</strong></td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td><strong>Missing values management</strong></td>
<td>Yes</td>
<td>Yes</td>
<td>Not discussed</td>
<td>Not standard</td>
</tr>
<tr>
<td><strong>Computational complexity of one run of the algorithm</strong></td>
<td>Linear in N</td>
<td>Quadratic in N</td>
<td>Linear in N</td>
<td>Linear in N</td>
</tr>
</tbody>
</table>

1. Step 1 or 2 stops converging.
2. If, for a specific cluster, the number of iterations necessary to decrease the relative difference between $R_k$ and $R_k_{PRELIM}$ (under ACCUR_RAD), is larger than a predefined number.
3. If the clusters removed from the data are not valid (number of genes below MIN_NR_GENES) found before one of the conditions of the stop criterion is true, is in practice also more or less proportional to the number of valid clusters found (e.g. for each invalid cluster found, two valid clusters will be found). This number of valid clusters is no classical attribute of the data (like $N$ or $E$) used to express computational complexity but it is rather a measure for the complexity of the structure of the data. Taken together, this means that the number of iterations in the global algorithms is also more or less proportional to this number of valid clusters in the data set and since the computational complexity of one iteration is approximately $O(N \times E)$, the computational complexity of the global algorithm is thus approximately $O(N \times E \times VC)$, ($VC =$ number of valid clusters). Notice also that, after finding a certain number of clusters, the number of genes left in the data is smaller than $N$ (clusters are discarded from the data). The computational complexity, as described above, is thus an upper limit.

**Computational and memory complexity of the global algorithm**

It is difficult to give an exact measure for the computational complexity of this heuristic approach. However, we can give an indication of the role of the most important variables. As previously said, the computational complexity of one cluster center localization is approximately $O(N \times E)$ ($N$ is the number of gene expression profiles in the data set, $E$ is the dimensionality of the expression vectors) and the computational complexity of the re-estimation of the cluster quality is negligible. So, the computational complexity of one iteration in the global algorithm (WHILE loop) is also approximately $O(N \times E)$. Notice also that Condition 2 of the stop criterion sets a limit for the maximum number of iterations in the global algorithm needed to define one cluster (which is only valid if the number of genes in this cluster equals or exceeds MIN_NR_GENES). Moreover, the number of invalid clusters (number of genes less than MIN_NR_GENES) found before one of the conditions of the stop criterion is true, is in practice also more or less proportional to the number of valid clusters found (e.g. for each invalid cluster found, two valid clusters will be found). This number of valid clusters is no classical attribute of the data (like $N$ or $E$) used to express computational complexity but it is rather a measure for the complexity of the structure of the data. Taken together, this means that the number of iterations in the global algorithms is also more or less proportional to this number of valid clusters in the data set and since the computational complexity of one iteration is approximately $O(N \times E)$, the computational complexity of the global algorithm is thus approximately $O(N \times E \times VC)$, ($VC =$ number of valid clusters). Notice also that, after finding a certain number of clusters, the number of genes left in the data is smaller than $N$ (clusters are discarded from the data). The computational complexity, as described above, is thus an upper limit.

Since only the distances from the expression profiles to the current cluster center have to be kept in memory (this is true at any stage of the algorithm), the memory complexity of the global algorithm is $O(N)$.

**RESULTS**

**Mitotic cell cycle of Saccharomyces cerevisiae**

The algorithm was tested on the expression profiling experiment of Cho et al. (1998), studying the yeast cell cycle in a synchronized culture on an Affymetrix chip...
(also see Spellman et al., 1998). This data set (http://cellcycle-www.stanford.edu) can be considered as a benchmark (Heyer et al., 1999; Jakt et al., 2001; Tamayo et al., 1999; Tavazoie et al., 1999; Yeung et al., 2001) and contains expression profiles for 6220 genes over 17 time points taken at 10-min intervals, covering nearly two full cell cycles. The majority of the genes included in the data set have been functionally classified (Mewes et al., 2000), which makes this data set an ideal candidate to correlate the results of new clustering algorithms with the biological reality. For more details about the data set itself, we refer to the original paper of Cho et al. (1998).

Our pre-processing included the following steps: data corresponding to the 90 and 100-min measurements were removed (Tavazoie et al., 1999). Also, we selected the 3000 most variable genes using $\sigma/\mu$ as a metric of variation (Tavazoie et al., 1999; filtering). Finally, we normalized the gene expression profiles as described in the normalization section. The results of the cluster analysis with our algorithm ($\text{MIN}_{NR}\_\text{GENES} = 10$, $S = 0.95$) are shown in Figure 3 (see supplementary web site). Table 1a summarizes the biological validation of this result by looking for enrichment of functional categories in individual clusters as described in Tavazoie et al. (1999). We mapped the genes in each cluster to the functional categories in the Munich Information Center for Protein Sequences (MIPS; Mewes et al., 2000) Comprehensive Yeast Genome Database. For each cluster we calculated $P$-values for observing the frequencies of genes in particular functional categories using the cumulative hypergeometric probability distribution. In the same table we also show, as a comparison and in parallel (where possible, we compare $P$-values of functionally matching clusters), the results obtained by Tavazoie et al. (1999) on the same data using the $K$-means algorithm. Note that the three most important clusters found by Tavazoie et al. (1999) (cluster 1, 4 and 2 in Tavazoie et al., 1999) could be matched with three clusters discovered by Adap_Cluster (cluster 1, 2 and 5). The degree of enrichment in the clusters identified by Adap_Cluster, however, was considerably higher and biologically more consistent.

In the biological validation and comparison discussed above, we filtered the data using the same metric of variance ($\sigma/\mu$) as proposed by Tavazoie et al. (1999) because different filtering strategies could produce different clusters independent of the clustering technique (we did not want different filtering to interfere with our comparison). However, in general, if filtering is performed, we recommend using simple measures of variation, like the standard deviation $\sigma$ (not $\sigma/\mu$) or the difference between the minimum and maximum value, together with Adap_Cluster. Using Adap_Cluster with the Cho et al. (1998) data indeed resulted in biologically more relevant results when using the standard deviation ($\sigma$) as the metric of variance. This analysis produced several clusters enriched in top-level functional categories (see Table 1b).

We were able to determine the role of every cluster presented in Table 1b within the yeast cell cycle context and correlate this role with the behaviour of the average profiles of the clusters. We have also found several protein complexes where nearly all members belong to the same cluster. A detailed discussion of these findings can be found on our supplementary web site.

Note that the results of Adap_Cluster in this section have been obtained without additional fine-tuning (we used the default value for $S$ of one or more parameters (unlike, for example, $K$-means; used by Tavazoie et al. (1999) where the number of clusters has to be estimated in advance, which is certainly not trivial) and that these results can be obtained very easily and almost instantaneously (maximum 1.5 min for the examples above on a typical PC).

**Additional results/simulations**

A discussion of the clusters found by our algorithm in other data sets (with different mathematical and biological characteristics) can be found on our supplementary web site:

- response to mechanical wounding in Arabidopsis (Reymond et al., 2000);
- central nervous system development (Wen et al., 1998);
- measurement of expression levels in different tissues (data not publicly available—manuscript in preparation);
- artificial data (with and without missing values).

**DISCUSSION**

The algorithm proposed in this paper is designed to find clusters of significantly coexpressed genes (higher degree of coexpression than could be expected by chance) in high-density areas of the data (high-density areas were assumed by Heyer et al. (1999) to be, biologically seen, the most interesting regions in the data). Genes not exhibiting an expression profile significantly similar to the expression profile of other genes in the data are not assigned to any of the clusters. The same applies to genes lying in low-density areas of the data. The size or radius for each cluster separately is determined—through the significance level $S$—by making a trade-off between the probability of false positive results (a gene assigned to the cluster that is not really coexpressed with the other members of the cluster) and the probability of false negative results (genes not assigned to the cluster.
but coexpressed with the members of the cluster). The default value for the significance level $S$ guarantees that a gene, which has been assigned to the cluster, has a probability of 95% or more to belong to the cluster (this means that the probability of being a false positive is 5% or less). In other words, the genes in the cluster are significantly coexpressed with a certain confidence. The significance level $S$, in turn, can be seen as a constant quality criterion for the clusters (while the quality criterion $R$ as defined in Equation (3) differs among the clusters defined by our algorithm). Our algorithm can thus be regarded as being a pure quality-based clustering method where all the clusters have a constant quality represented by the significance level $S$ (the term adaptive quality-based clustering is thus only valid when using Equation (3) as quality criterion). When compared to the previous definition (quality measure $R$), this new quality measure $S$ has the advantage that it has a strict statistical meaning (it is much less arbitrary) and that, in most cases, it can be chosen independently of a specific data set or cluster. In addition, it allows for the setting of a meaningful default value (95%).

In Table 2a a detailed comparison between our global algorithm (Adap_Cluster) and the algorithm proposed by Heyer et al. (1999; QT_Clust) is made. Because we focus on algorithmic aspects, the QT_Clust algorithm in our comparison uses the same distance and quality measure as we did (Euclidean distance and quality defined as in Equation (3)—In Heyer et al. (1999) the jackknife correlation was used together with a quality measure defined as a diameter). This change of distance and quality measure does not significantly change the structure of QT_Clust and in essence, there is no fundamental difference between a quality defined as a radius and a quality defined as a diameter.

To complete the picture, Table 2b gives a summary of the differences between our method, hierarchical clustering, SOM and $K$-means.

Clusters formed by our algorithm might be good ‘seeds’ for further analysis of expression data (Thijs et al., 2002) since they only contain a limited number of false positives. When the presence of false positives in a cluster is undesirable, a more stringent value for the significance level $S$ might be applied (e.g. 99%; for noise-sensitive analyses such as motif finding) which will result in smaller clusters exhibiting a more tightly related expression profile.

Conclusively, Adap_Cluster can be considered as an intuitively appealing, user-friendly (no need for a predefinition of the number of clusters, statistical and intuitive definition of the quality measure with a meaningful default value) and fast clustering algorithm.

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REFERENCES


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