The current excitement in bioinformatics – [Au: OK?] analysis of whole-genome expression data: how does it relate to protein structure and function?
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Whole-genome expression profiles provide a rich new data-trove for bioinformatics. Initial analyses of the profiles have included clustering and cross-referencing to ‘external’ information on protein structure and function. Expression profile clusters do relate to protein function, but the correlation is not perfect, with the discrepancies partially resulting from the difficulty in consistently defining function. Other attributes of proteins can also be related to expression — in particular, structure and localization — and sometimes show a clearer relationship than function.

Introduction
Bioinformatics has traditionally involved the computational analysis of large molecular biology datasets. Initially, these were drawn from the world of protein structure. In 1995, the field changed with the advent of complete genome sequences, which represented a new type of large-scale data. Now, whole-genome expression experiments are providing further sources of large-scale data and transforming bioinformatics yet again. Expression experiments can generate a quantity of information that potentially dwarfs that provided by [Au: OK?] genome sequences and protein structures. Whereas it is sufficient, for many practical purposes, to view genome sequencing as a one-time process for each organism (except for the analysis of individual genetic variations), expression experiments can be repeated an arbitrary number of times to monitor the expression of different cell types and states (diseased or healthy), or the same cells at different times or in different individuals. The number of potential experiments is only limited by cost and imagination. Each of these experiments potentially gives rise to a new genome-scale dataset and a further challenge for bioinformaticians.

Expression data
Technologies and systems: SAGE, chips and arrays in yeast and beyond
Genome-wide expression information is principally generated by three technologies: cDNA microarrays [1], GeneChips (also called high-density oligonucleotide arrays) [2] and SAGE (serial analysis of gene expression) [3]. These technologies, which are all new and rapidly evolving, have been recently reviewed [4–6]. The large number of ESTs (expressed sequence tags) [Au: OK?] in different cells and tissues provides a further source of large-scale expression information [7].

Expression monitoring on a genome-wide scale was first successfully demonstrated in yeast [8–10]. Later experiments have been performed on other organisms, including mycobacteria [11], Escherichia coli [12], worm (see http://bioinfo.mbb.yale.edu/genome/expression [Au: OK?]), fly [13], mouse [14] and human [15,16]. There are a number of technical difficulties associated with certain systems (e.g. the lack of poly-A tails in bacteria) but, in principle, these experiments can be applied repeatedly in a wide variety of organisms.

Relevant for computations: absolute versus relative, population averages and [Au: OK?] databases
From a computational perspective, the three expression technologies all produce a profile (or vector) of expression levels for many genes. In principle, GeneChips and SAGE allow the measurement of absolute expression levels (in units of mRNA transcripts per cell), whereas cDNA microarrays primarily measure changes relative to a reference state (yielding an ‘expression ratio’). Although valuable, absolute transcript abundance measurements do not completely measure mRNA concentration, which also depends on cellular compartment volume.

Expression experiments measure cell population averages, not individual cells, so another important issue is the degree to which all cells in the investigated population are in the same ‘state’. For single-cell organisms, temporal synchronization can often be achieved artificially, for example, [Au: OK?] in yeast cell-cycle experiments, cyclins were used for synchronization [10,17]. Work in multicellular organisms has the added complexity that expression measurements may combine many different tissues. Recent papers have discussed statistical aspects of expression data in detail [18,19*].

The first major bioinformatics task related to expression data is organization and storage. This is currently the sub-
ject of much discussion and there are a number of pilot databases: GEO (the NCBI Gene Expression Omnibus); ExpressDB [20] (Harvard); GeneX (NCGR); the Stanford Microarray Database; and ArrayExpress [21] (see http://bioinfo.mbb.yale.edu/genome/expression [Au: Does this link refer to all the databases or just ArrayExpress?]). Some issues being considered include [Au: OK?] whether to normalize and standardize the data, whether it should be stored in a central archive or federation of web sites and to what degree details about experimental design should be kept. Storing the raw array intensities lends itself nicely to standard relational tables. However, information related to the experimental conditions (tissues, drug treatments, etc.) is more complicated. To some degree, how to best archive the data will be determined by the most popular analyses that bioinformaticians end up performing.

**Computational issues: internal versus external, supervised versus unsupervised**

Analysis of expression datasets [Au: OK?] encourages more exploratory, data-driven styles of research than traditional
Figure 1 legend

An overview of some principles of expression data analysis. The top part of the figure shows a representation of the input data. Expression data consist of expression level measurements for various genes arranged in ‘profiles’ across either different conditions or different times. One can determine the distance between each pair of profiles and put this into a large ‘distance matrix’, which then forms the basis for many of the clustering algorithms. (This is also known as a ‘correlation matrix’ or ‘kernel matrix’ in various calculations.) The top part also gives a schematization of the types of external information expression profiles can be related to. It shows part of a genome with the relationship between transcribed gene sequences and protein structure and function. Note how a number of genes can share the same protein fold, how certain protein folds can have many functions and how two different folds can have the same function. The bottom part of the figure illustrates a number of ways of analyzing expression data. Broadly, these can be divided into calculations dealing purely with the internal structure of the profiles and calculations relating the profiles to external, nonexpression information. Specific examples of various methods for analyzing the correlation matrix of expression profiles are PCA [34••], k-means clustering [30••] and SOMs [31••].

Hierarchical clustering [22••,29•] and SVMs [24••]. PCA tries to find the directions of greatest variance implied by the correlation matrix and to then ‘visualize’ the data in terms of their projection on these directions. Hierarchical clustering successively groups together the profiles that are the most similar, generating a tree-like description of the data. There are a variety of ways of making this determination of similarity; for example, in UPGMA, it is based on the distance to an existing averaged group center, whereas in single-linkage, it is based only on the distance to the nearest representative of a given cluster. K-means clustering algorithms make few assumptions about the data. They start with a number (k) of randomly positioned cluster centers and then update their positions to fit the data. SOMs are similar, but they impose a bit more structure on the clustering, requiring that the updated position of a cluster be affected by the position of the other cluster centers. In relation to the subschematic illustrating SOMs, adapted from [31••], note that SOMs would have constraints related to the dotted lines, whereas in k-means these would be absent. SVMs assume that the profiles are ‘tagged’ with already known classification information, such as a functional class. They then implicitly transform the data into a higher dimensional representation in which a simple plane can be found to separate the differently tagged groups. (In practice, this is accomplished by considering nonlinear measures for distance, beyond simple correlation.) SVMs are considered a type of supervised learning, in that they explicitly train and test against the external data. In contrast, SOMs, k-means and hierarchical clustering are considered unsupervised clustering, in that they do not relate the learned clusters to the external data until after they have been derived. However, one could imagine various unsupervised algorithms that simultaneously consider expression data and additional features derived from external information (such as localization) in learning clusters.

Clustering: bottom-up hierarchies versus top-down partitions

The main type of internal analyses involves clustering and partitioning the data. As schematized in Figure 1, the starting point for clustering methods is defining a similarity measure among expression profiles and then constructing a matrix giving a distance between each pair of profiles. In general, there are many possible metrics [23••,24••]. A common one is the Pearson correlation coefficient [22••,25••]. An interesting modification of this is the ‘jackknife correlation’, which is robust with respect to data outliers [26].

Hierarchical methods group profiles in a ‘bottom-up’ fashion, joining the most similar profiles into clusters first and then including more diverse ones [27]. There are a variety of specific approaches (e.g. UPGMA [Au: Would it be helpful to briefly explain what this is?], single-linkage, multiple-linkage, etc.), which were mostly derived from phylogenetic tree construction [28]. These were the first methods applied to expression data [22••,25••,29] and they have the advantage that the number of clusters needs not be specified beforehand. However, their drawback is that there is no reason to believe that expression data — in contrast to evolutionary information — is naturally organized in bifurcating trees. The trees produced by hierarchical clustering can only be broken into clusters in some ad hoc fashion. Furthermore, decisions made early in bottom-up clustering cannot be undone and sometimes adversely affect the final result.

In contrast to bottom-up clustering, partitioning approaches are ‘top down’. Important examples applied to expression analysis are k-means [30••] and self-organizing maps (SOMs) [31••,32•]. A tree structure is not assumed in these methods; however, they often require an a priori decision on the number and structure of distinct clusters. It remains a problem to objectively determine the optimum number of clusters for these algorithms [19]. Recently, partitioning algorithms have been developed in
which the number of clusters is determined by the algorithm itself [33•].

An additional method of internal analysis is principal component analysis (PCA) [34•]. This method can be understood [Au: Would ‘used’ perhaps be better here?] as a way of compressing the data and filtering out noise by projection onto a low-dimensional subspace. It can be used for data visualization and initial exploration of clusters.

Phenotype characterization: cancer diagnosis

Another type of internal analysis uses [Au: OK?] expression patterns to distinguish between cell types and disease states. In this context, entire expression profiles can be used to compare different experiments [Au: Use inverted commas?] (in contrast to clustering genes). There have already been many applications in cancer diagnosis [35••,36,37•]; however, a full discussion is beyond the scope of this review.

Relating expression profiles to protein function

Thus far, we have only discussed computations aimed at revealing the internal structure of [Au: OK?] expression data. Expert biological knowledge is applied afterwards to interpret the results. The next type of analysis tries to explicitly integrate information about protein function, structure and so forth directly into the expression data computations. First, we will look at work relating expression profiles to protein function. As a preamble, it is worthwhile to briefly discuss how protein functions are classified.

Functional classification and its problems

There are a number of schemes for classifying protein function, which have been recently reviewed [38]. Briefly, most of the schemes concentrate on a single organism, for example, MIPS for yeast, GenProtEC for E. coli, FlyBase for Drosophila and EGAD for human ESTs [39,40] (see http://bioinfo.mbb.yale.edu/genome/expression). Other schemes classify a subset of functions across a variety of organisms, for example, ENZYME for enzyme function and EcoCyc, WIT and KEGG for pathways [41–44]. There

\[ R_{ij} = \frac{1}{N-1} \sum_{k=1}^{N} x_{ik} \cdot x_{jk} \]

where \( N \) is the number of elements in the profiles \( x_i \) and \( x_j \). The normalized profile \( x \) can be computed as a ‘Z-score’ from the measured expression ratio profile \( x \) through the relation

\[ x(k) = \frac{x(k) - x_{avg}}{\sigma} \]

where \( x_{avg} \) denotes the average and \( \sigma \) the standard deviation of values in \( x \), and \( x(k) \) and \( x(k) \) are the \( k \)th components of their respective profiles. Given a group of genes, we can compute the correlation coefficient matrix \( R \), where each element \( (R) \) of the matrix denotes the Pearson correlation coefficient between genes \( i \) and \( j \). We can then compute an average correlation coefficient \( R_{avg} \) by averaging the matrix elements (excluding the main diagonal). This statistic gives an idea of the overall similarity of the expression profiles in a group of genes. Although there are \( O(G^2) \) elements in \( R \), the computation time for \( R_{avg} \) can be kept proportional to \( O(G) \) by calculating \( R_{avg} \) as follows:

\[ R_{avg} = \frac{1}{G^2 - G} \sum_{i=0}^{G} \sum_{j=0}^{G} R_{ij} - G \]

where

\[ X_{Tot} = \sum_{g=1}^{G} X_g \]

is the sum of all expression profiles in the group of genes. The figure shows the distribution of this statistic for the expression data measured during the diauxic shift in yeast [8]. Groups of genes of size \( G \) were randomly chosen from the genome. For \( G = 2 \), the statistic is simply the Pearson correlation coefficient itself. For increasing \( G \), the distributions become narrower. The distributions were generated by sampling \( R_{avg} \) 10,000 times from the full distance matrix relating the expression profiles of all approximately 6000 genes in yeast. Functions of the form

\[ F(y) = \sum_{i=0}^{6} a_i y^i \]

can be fit to the cumulative distributions, where

\[ y = \ln \left( \frac{2}{1 - R_{avg}} \right) \]

is a transformation of the average correlation coefficient \( R_{avg} \) with \( a_0 = b_2 = 1, a_2 = a_0 = 0 \). For the graph shown in the figure (\( G = 2 \)), we used parameters \( a_0 = 19.92, a_2 = 5.66, a_1 = -3.22, b_2 = -1.02, b_0 = 2.07, b_1 = 28.97, b_3 = 3.47, b_4 = 4.56, b_5 = -1.56, b_6 = -1.21 \).
have been [Au: OK?] some attempts to merge functional classifications for different organisms into one common source (the Gene Ontology Project [45], see http://bioinfo.mbb.yale.edu/genome/expression), although the creation of a complete universal functional system will be a difficult task [38,46]. However, there have been some attempts in terms of creating unique keyword combinations or sequence variability signatures for functions [47,48].

Beyond the lack of scope of the current classification schemes, it is important to realize that there are many profound difficulties in functional classification. First, the concept of ‘function’ is itself rather vague. Sometimes it is defined in terms of biochemical mechanism (e.g. ‘adenylate kinase’); at other times, in terms of either [Au: OK?] involvement in pathways or overall cellular role (e.g. part of ‘glycolysis’ or ‘cellular metabolism’); and, finally, sometimes in terms of the phenotype of the organism when the associated gene is disabled (e.g. ‘causes cancer’). [Au: I

![Figure 3](image-url)

Current Opinion in Structural Biology
Figure 3 legend

The degree of expression profile similarity is different for genes from different functional groups and also varies between different expression experiments. We illustrate this concept in the context of the MIPS functional classification scheme. Each part shows the negative logarithm of the one-sided P-values (–log(P)) based on distributions of the average correlation coefficient for different experiments, as explained in the legend to Figure 2. The P-values give the probability that an average correlation greater than that observed for each functional group could have arisen from a randomly selected group of genes of the same size. Accordingly, lower P-values or higher values of –log(P) indicate a greater significance of the similarity between expression profiles. The P-values range from 0 to 1; correspondingly, –log(P) ranges from infinity to 0. For values of –log(P) greater than four, we cannot determine the value with certainty because of the limited scale of our computation; we indicate this in the table by ‘>4’ (for highly significant groupings). [Au: What does the shading in the table represent? The degree of significance of the grouping. Please explain.] Each row in parts (a) and (b) of the figure corresponds to an MIPS functional category and each column corresponds to a different expression experiment on yeast. The first experiment is a GeneChip experiment [17] to monitor the cell cycle synchronized by the cyclin CDC28. The other experiments are microarray experiments: the cell cycle synchronized by CDC15 [10], the diauxic shift [9] and the process of sporulation [8]. Part (a) shows the most general MIPS categories, whereas (b) shows the subcategories of the top-level MIPS category ‘energy’.

Part (c) summarizes the fraction of functional categories that represent ‘significant’ groupings with respect to expression. We define a grouping as significant if we find values of –log(P) > 3, a less than 1 in 1000 chance that the observed average correlation arises randomly. The first column indicates the level in the MIPS hierarchy. (MIPS 1 is the first level, MIPS 2 is the second level, etc.) The next columns show the fraction of significant groups for each experiment and the last column shows the total number of groups in each MIPS level. The fraction of significant groups decreases as the detail of classification increases from the first to the third MIPS level. This is because (for the quantitative assessment presented here) a high significance for a more specialized MIPS category tends to also show up in a high significance for the more general MIPS category one level above. In part (c), we show the significance of the clustering determined by various methods described in the text – in particular, hierarchical clustering [21], k-means [30×] and SOMs [2]. The hierarchical clustering was applied to all four experiments and to additional data on the mitotic cell cycle, and temperature and reducing shocks (see http://bioinfo.mbb.yale.edu genome/expression). To apply the methodology, the hierarchical tree was cut off such that 25 ‘subtrees’ or gene clusters remained. Clearly, both these methods produce much more statistically significant clustering with respect to expression than the MIPS functional groups. The only functional categories for which we find high significance in all four experiments are at the top of the table: ‘cell growth, division and DNA synthesis’ and ‘protein synthesis’ (including ribosomal proteins). In contrast, some categories are not significant in any of the experiments (such as ‘beta-oxidation of fatty acids’, another subcategory of ‘energy’). In general, there seems to be a higher degree of correlation for the two cell-cycle experiments than for the other two experiments (e.g. for the ‘metabolism’ category), perhaps because the mechanics of the cell cycle forces a high degree of transcriptional coexpression on many functional systems. However, a few functional groups show a higher significance in the diauxic shift and sporulation experiments (such as the group ‘glyoxylate cycle’, which is a subcategory of ‘energy’). It can be clearly seen that many functional groups show different degrees of coexpression under different experimental conditions, highlighting the importance of experimental design.

have bracketed the examples in order to make a long sentence more digestible. Please let me know if you think that I have failed!] Second, many proteins are multifunctional, having more than one function, sometimes in unrelated areas [49]. For instance, the protease thrombin is primarily associated with blood clotting, but also interacts with receptors for cell activation and neural development [50]. Third, conversely, multiple gene products often collectively carry out a single function (e.g. the ribosome). Fourth, the naming of functions is currently unsystematic and inappropriate for quantitative comparisons. Humorous examples of this come from the fly, for which [Au: OK?] genes have most bizarre names, for example, ‘suppressor-of-white-apricot’ and ‘darkener-of-apricot’, which are, respectively, an RNA-binding protein and a kinase involved in eye-color determination (SUWA_DROME and DOA_DROME [Au: Please explain what these terms relate to]). There have been some attempts in terms of creating unique keyword combinations or objective sequence variability signatures for functions [47,51,52].

Supervised learning (support vector machines)

Given a function classification, one would like to know how well clusters of expression profiles relate to functional categories and, if there is a relation, the degree to which it can be used to predict the functions of genes. Some initial reports on expression analysis suggested that certain prominent expression clusters did relate to functional categories and that function prediction was possible [22×,53,54]. More recent work has tried to systematically test this proposition using explicit training and testing sets. As diagramed in Figure 1, one technique that has been applied is support vector machines (SVMs) [24×]. This supervised learning technique positions a hyperplane to partition the data and minimize the number of misclassified proteins on the basis of a known functional classification or empirical measurements not included in the dataset.

Other supervised learning approaches include decision trees, Parzen windows and Fisher’s linear discriminant [24×]. More general approaches that make use of prior information include Bayesian networks [55].

Global characterization of the expression/function relationship

The calculations relating expression and function have largely focused on specific cases or functional categories. Figures 2 and 3 attempt to give an overview of how they relate in a ‘global’ sense. On the basis of [Au: OK?] the results of a whole-genome expression experiment, one can determine the distribution of similarity values for each pair of genes, that is, the distribution of the [Au: OK?] correlations of their expression profiles. For groupings larger than
pairs (e.g. triplets), this can be generalized to the distribution of the average value of the correlation. Sample distributions based on the yeast [Au: OK?] diauxic shift timecourse [9] are shown in Figure 2. And, as shown in Figure 3, with respect to any particular expression experiment, distributions can be used to evaluate the statistical significance of a given clustering of genes. Most of the clusters automatically generated using the algorithms discussed earlier (e.g. hierarchical clusters or SOMs) appear to be significant. For instance, on the basis of a P < .001 threshold, 28 of the 30 SOM clusters for the cell-cycle data are significant (93%). However, fewer gene groupings based on the functional categories in MIPS are significant, for example, only 10 of the 16 top-level MIPS clusters have P < .001 (63%) for the same experiment. Some functional groups are always highly correlated with expression profiles (e.g. ‘cell growth’ and ‘protein synthesis’). However, other MIPS groups are only correlated in certain experimen-t
This figure shows how expression data can be related to protein structure. It shows a number of protein folds in the yeast, E. coli and C. elegans genomes ranked by various measures related to expression [Au: Again, what does the different shading of the rankings show?]. At least the four most common folds for each type of ranking are shown. The first column shows the rank of the fold in terms of how many times it is found in the yeast genome (i.e. by duplication), based on recent PSI-BLAST structural assignments [63]. The next column shows its ranking in the transcriptome [64•, 65•], that is, the occurrence of each fold weighted by the number of copies of mRNA associated with it, based on GeneChip data [75]. Folds can be also ranked in terms of their fluctuation in mRNA levels over an experiment, rather than their total number of mRNA copies, using the average standard deviation of the expression ratios as an indication of the degree of fluctuation. Such rankings are shown in the next four columns. Columns 3 and 4 show a ranking based on the fluctuation in expression in the yeast cell cycle (CDC28 [17] and CDC15 [10]). Columns 5 and 6 show rankings based on other yeast experiments, the diauxic shift [9] and sporulation [8]. For comparison, columns 7 and 8 show the ranking for other organisms, E. coli (based on fluctuation in the heat shock experiment [12]) and C. elegans (based on the fluctuations during successive larval stages of the worm; V Reinke, personal communication). [Au: Does column 8 not give the PDB codes of the proteins, rather than the C. elegans ranking? Have I got the correct figure? Please can you check and adjust the legend accordingly.] Note how different all the rankings are. The most common folds in the transcriptome have a mixed coil/structural architecture and are mostly cytosolic enzymes. The most abundant fold is the TIM barrel, which is also known to be the most versatile fold, associated with 16 different enzymatic functions [83]. In terms of the fluctuation rankings, one fold that changes considerably in expression is that of ATPase domain of HSP90/DNA topoisomerase II, which is highly ranked in both cell-cycle experiments (CDC28 and CDC15) and the E. coli experiment. The folds are selected from the current 520 folds and 771 superfamilies as of 1 November 1999 in SCOP 1.48 [56]. For the yeast fluctuation rankings, we excluded genes with an absolute expression level lower than 100 units of intensity, as given by the CDC28 GeneChip, because the signal fluctuations of lowly expressed genes are most likely due to measurement uncertainties. (The absolute expression level is defined as the difference between the intensity of the oligonucleotide-perfect match [PM] and the background intensity measured by a single mismatch probe [MM].). For the E. coli experiment, we simply ranked the expression ratio because no time series measurements were taken. For the C. elegans fluctuation ranking, we excluded signals with less than 250,000 units [Au: Again, is there a C. elegans ranking in the figure?]. [Au: What is ‘dim dom’ with respect to amino acid dehydrogenase? What is TBP? What is shown by the Rep. PDB column?]
Relating expression data to other external information

Another attribute of a gene that can be related to its expression profile is its regulation. This subject has been reviewed in detail [66], so we will only touch upon it briefly here. Almost by definition, genes that have similar expression profiles probably share upstream regulatory elements. This fact has been exploited to search for new regulatory sequences [30••,67–69]. For genes that have similar expression profiles but do not share an obvious regulatory element, one can use an unsupervised motif learner, such as a Gibbs sampler [70], to discover new regulatory motifs in upstream sequences.

Other attributes of proteins that have been related to expression include subcellular localization and protein–protein interactions. As was the case with protein structure, these attributes of proteins can be more precisely systematized than function. For yeast, systematic information on localization and interactions is tabulated in the MIPS, YPD and SwissProt databases [40,71,72]. With regard to localization, it has been found that cytosolic proteins tend to be expressed at high levels, whereas proteins destined for membranes and mitochondria are expressed at lower levels [73•]. Proteins in the secretory pathway have high fluctuations in expression level over timecourses. Collectively, this information can, in fact, be combined to help predict the localization of proteins for which there is expression information available, but no known localization [74].

Conclusions

The advent of whole-genome expression experiments has led to a new class of bioinformatics analyses. These fall into two main groups: internal clustering and comparison of expression data, and cross-referencing of expression data to other information on protein structure and function. With respect to the experiments on yeast, clusters of genes that have similar expression profiles often fall into the same functional category. However, this is not always true in a ‘global’ sense. The discrepancies reflect particular functional categories highlighted by certain experiments. More importantly, they also result from the difficulty in consistently defining function across a wide variety of proteins. We believe this latter difficulty is quite significant and probably the major current impediment to interpreting expression data in terms of protein function. We can side-step this to some degree by focusing on attributes of proteins other than function, such as structure, regulation and localization. Many of these can be defined in a much more consistent fashion than function and, perhaps because of this, show a clearer relation to gene expression.

Acknowledgements

On the web, we will make available supplementary data related to the review (extended versions of Figures 2, 3 and 4, with a list of fold expression levels and function significance values for the whole yeast genome) and a ‘links page’ to web sites referred to in the text. Go to http://bioinfo.mbb.yale.edu/genome/expression [Au: This is probably the best place to put this (there is no precedent for this sort of supplementary material in the Current Opinion journal). It is a very good idea.]

References and recommended reading

Papers of particular interest, published within the annual period of review, have been highlighted as:

* of special interest
** of outstanding interest


The authors demonstrate the application of self-organizing maps (SOMs) to cluster gene expression data of the yeast cell cycle and hematopoietic differentiation. An advantage of the method is the short computation time. SOMs are one method of extracting the most prominent patterns from a dataset. They are similar to k-means, but assume a more structured (correlated) distribution of cluster centers. Although this approach does appear to work well on expression profiles, it is not clear what in expression data justifies the added structure in the SOM approach.


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Independently of the work described in [31*], the authors demonstrate the application of self-organizing maps to the expression data for the diauxic shift [9]. The self-organizing maps algorithm is an unsupervised neural network learning algorithm.


The authors describe the theoretical and practical application of a partitioning clustering algorithm that recovers an original cluster structure with high probability when the expression data exhibits stochastic errors.

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[Au: Each bulleted reference must be accompanied by an annotation. Please either delete the bullet or provide a few sentences describing the results of the paper and their implications.]


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74. Drawd A, Jansen R, Gerstein M: Gene expression levels are related to protein subcellular localization. Trends Genet 2000, in press. [Au: Do you have any more details (volume and page range) for this paper?] It was observed that highly expressed proteins tend to be cytosolic, whereas lightly expressed ones tend to be localized in the nucleus or the membranes. The relation between a gene’s expression level and the subcellular localization of its associated protein is then used to help predict the localization of the more than 4000 yeast proteins with unknown localization. A probabilistic Bayesian formalism is used, whereby the localization of a protein is iteratively updated depending on a variety of features it has. These features include expression level and also the presence of [Au: OK?] traditional sequence motifs (e.g. HDEL).