

Research

Strong-association-rule mining for large-scale gene-expression data analysis: a case study on human SAGE data

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Abstract

Background: The association-rules discovery (ARD) technique has yet to be applied to gene-expression data analysis. Even in the absence of previous biological knowledge, it should identify sets of genes whose expression is correlated. The first association-rule miners appeared six years ago and proved efficient at dealing with sparse and weakly correlated data. A huge international research effort has led to new algorithms for tackling difficult contexts and these are particularly suited to analysis of large gene-expression matrices. To validate the ARD technique we have applied it to freely available human serial analysis of gene expression (SAGE) data.

Results: The approach described here enables us to designate sets of strong association rules. We normalized the SAGE data before applying our association rule miner. Depending on the discretization algorithm used, different properties of the data were highlighted. Both common and specific interpretations could be made from the extracted rules. In each and every case the extracted collections of rules indicated that a very strong co-regulation of mRNA encoding ribosomal proteins occurs in the dataset. Several rules associating proteins involved in signal transduction were obtained and analyzed, some pointing to yet-unexplored directions. Furthermore, by examining a subset of these rules, we were able both to reassign a wrongly labeled tag, and to propose a function for an expressed sequence tag encoding a protein of unknown function.

Conclusions: We show that ARD is a promising technique that turns out to be complementary to existing gene-expression clustering techniques.

Background

We are now entering the post-genome era and it seems obvious that, in a near future, the critical need will not be to generate data, but to derive knowledge from huge datasets

generated at very high throughput. This has been a challenge for quite some time in genomic research, and is now a challenge for transcriptome research, that is, the analysis of gene-expression data.

The kind of raw data in which we are interested can be considered as a matrix, denoted as G , of real numbers (Table 1). The rows denote different samples or conditions, such as the same cell type in different biological situations, and are indicated in this hypothetical example by Greek letters. The columns, indicated by letters, denote genes. In practice, we can have hundreds of lines and thousands of columns. $G[\alpha,c]$ denotes the quantitative expression of gene c in the situation α . Table 1 shows a model matrix that we will use in our explanations later.

Most of the available gene-expression data-analysis methods are based on clustering algorithms that try to establish syn-expression groups [1], that is, groups of genes whose expression is correlated in different biological situations. The basis for all clustering algorithms is their ability to generate groups of genes that fulfill two related constraints: intra-group similarities should be maximized and intergroup similarities should be minimized.

Although such algorithms have been quite successful, most notably in the molecular profiling of human cancers [2], their biological validity can be questioned when the identification of molecular networks is the goal. In this context, they have three main drawbacks. First, a gene which functions in numerous physiological pathways, such as that for the p53 protein [3], will have to be clustered in one and only one group. Second, no relationship can be inferred between the different members of a group. That is, a gene and its target genes will be co-clustered, but the type of relationship cannot be rendered explicit by the algorithm. Third, most clustering algorithms will make comparisons between the gene-expression patterns in all the conditions examined. They will therefore miss a gene grouping that only arises in a subset of cells or conditions.

To overcome these problems, we investigated the potential impact of the association-rule discovery (ARD) technique.

Table 1

Example matrix for gene-expression data

	a	b	c	d
α	22	73	57	17
β	57	32	64	21
γ	41	83	21	23
δ	65	81	59	70
ϵ	18	79	56	64
ρ	52	39	59	16
π	30	76	61	65

The rows denote different samples or conditions, such as the same cell type in different biological situations while the columns denote genes. $G[\gamma,c]$ denotes the quantitative expression of gene c in the situation γ .

This is an unsupervised data-mining technique that seeks descriptive rules in potentially very large datasets [4]. This method should resolve the above drawbacks of existing clustering approaches for the following reasons. First, any gene can be assigned to any number of rules as long as its expression fulfills the assignment criteria. This means that a gene involved in many synexpression groups will appear in each and every one of those groups, without limitation. Second, rules are orientated (If ... then ...) and thus to a certain extent describe the direction of a relationship. For example, a gene encoding a transcription factor should appear in the left portion of the rule and its target genes in the right portion. Third, by exploring low values of frequency, one can identify rules that are true in only a limited number of cells or situations. This means that if, in the overall dataset, a specific subset of cells exhibit highly characteristic patterns of gene expression, the algorithm should be able to detect it. Last but not least, by focusing on strong rules, the biologist does not have to browse and study a huge number of redundant rules.

Contribution

In this paper we evaluate the ARD for generating synexpression groups from large gene-expression matrices. The kind of rule we wish to discover is, for example, ‘When gene a and gene b are overexpressed within a situation, then often gene c is overexpressed too’. Such a rule will be designated valid. If it is also discovered that genes a , b and c are often overexpressed in various situations, then the rule would be designated frequent. The user simply defines thresholds for ‘frequent enough’ and ‘valid enough’, but no other information is needed. An association-rule miner then identifies every rule that is frequent and valid according to these user-defined thresholds. This is clearly unsupervised data mining.

This process is quite complex, especially when the number of columns is very large, which is typical of gene-expression data. Indeed, the number of potential rules increases exponentially with respect to the number of columns, and it rapidly becomes impossible to enumerate these rules and check if they are frequent and valid. As well as this aspect of complexity, it is also necessary to consider that not all the frequent and valid rules may be interesting to the biologist. A second processing phase will be required to identify the most promising rules first. Here again, however, we assume that this concept of ‘promising’ cannot be determined in advance.

We assessed the potential of the ARD technique by using the Min-Ex approach (see below), which provides an elegant solution for eliminating redundant association rules. This has been recognized as a major improvement in the standard association-rule mining technique [5]. As this improvement should greatly facilitate the extraction of interesting rules from very large, densely correlated and highly dimensional data, we used ARD with Min-Ex for analyzing publicly available data

obtained by serial analysis of gene expression (SAGE) in human cells [6,7].

Results

SAGE is an experimental technique designed to quantify gene expression [8]. This technique produces 10-base sequences, or ‘tags’, that identify one (or more) mRNAs. The frequency of each tag in the overall produced sequence reflects the cellular abundance of the corresponding transcript(s).

SAGE data have been poorly exploited by data-mining techniques so far; we are aware of only one study on the complete set of data [9]. One obvious reason for such poor exploitation lies in the structure of the data, including a high error rate for low-frequency tags (and especially tags appearing only once in a library), and a very high dimensionality.

The use of different approaches for generating the boolean values (see below) that indicate whether a gene is overexpressed or not solves the problem of low-frequency tags. Furthermore, the Min-Ex algorithm that is implemented in the ac-miner software we used has been designed for handling huge and highly dimensional datasets.

We explored the validity of the association-rule technique after generating different types of boolean matrices and comparing the rules extracted in each case. We also compared the data we obtained with those obtained through ‘traditional’ clustering, and conclude that clustering and association rules can be seen as complementary.

Introducing the raw data

The SAGE data were downloaded through ftp [10] as of June 2001. We downloaded: the tags_freq files (list of unique tags with frequency seen [11]), the description files [12] and a

Stats.txt spreadsheet file [13] containing various information regarding the libraries, and establishing a link between the tags_freq files and the description files.

The minimal transcriptome set was generously provided as an Excel file by V. E. Velculescu (Johns Hopkins Oncology Center, Baltimore). This consisted of 1,183 tags, their expression values in different tissues and their identification.

Preprocessing raw data and preparing the boolean context

We used only the human SAGE libraries that were found in the Stat.txt file. This left us with 93 libraries. So that we would not have to analyze the situation created by missing data we limited ourselves to the genes belonging to the minimal transcriptome set [14]. We understood this set as being genes that should be found in each and every human library analyzed. This was not the case, and the vast majority of those tags were not found in one or other library. As this might be due to the very small size of some libraries (Figure 1), we tested the percentage of missing tags as a function of library size. The smaller libraries (less than 20,000 tags) indeed harbored a very high percentage of missing tags. Using a cutoff value of 20,000 sequenced tags, we obtained a frequency of missing tags of 8%. This value was not considerably modified when a much higher cutoff value (50,000 tags, 5% missing tags) was applied; this value, however, resulted in a great loss of information. We therefore decided to use the 74 libraries that displayed more than 20,000 tags, and we decided to give a value of zero when a given tag was missing in a given library. As our approach is based on the use of a cutoff value (see below) we feel quite confident that this should have only a marginal effect, if any, on the output. Furthermore, the resulting expression profiles did not display any aberrant form due to the use of the null value (see below).

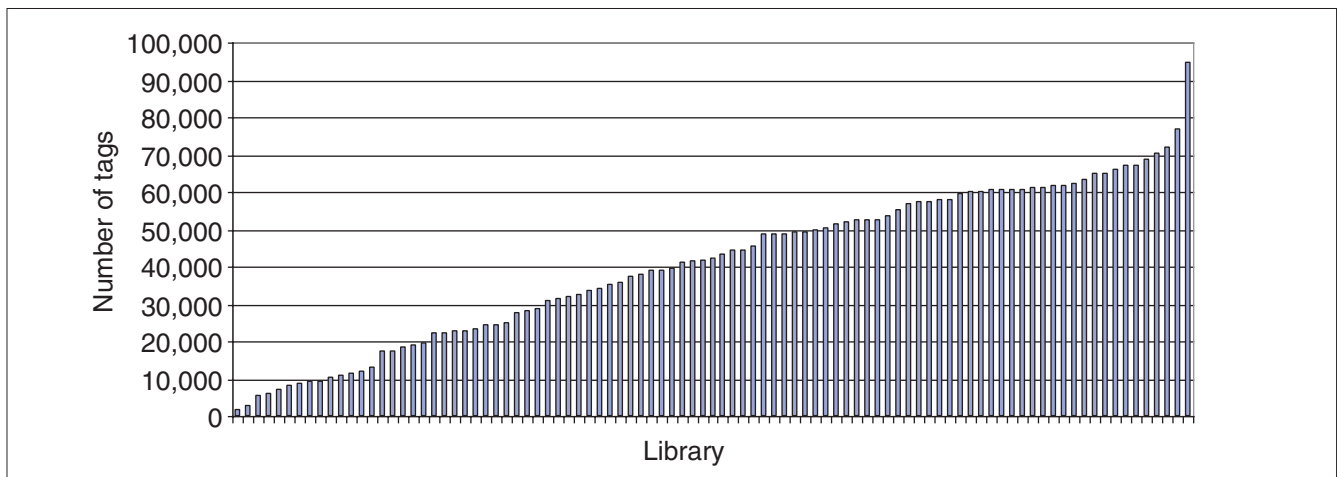


Figure 1
Size distribution among the 93 SAGE human libraries, in overall number of sequenced tags.

Finally, we re-established a strictly minimal transcriptome set containing tags found in each and every library with more than 20,000 tags (Figure 2). This set contained 156 tags, among which we found 56 representing genes encoding ribosomal proteins, and 15 representing genes involved in translation processes. Of the original 1,183 tags belonging to the minimal transcriptome, 361 were found to represent ambiguous tags (a single tag sequence matching more than one gene description) in the original Velculescu Excel file. Those 361 ambiguous tags were discarded. We therefore continued with a list of 822 non-ambiguous tags.

The original tags_freq files contained the following information: a list of tag sequences, and the number of times each sequence was found in the library. Because the actual library size is highly variable (Figure 1), such sequences could not be compared without normalization. We used the same normalization approach as previously described [9]: the tags_freq value was divided by the total number of tags in the library and multiplied by 300,000, the estimated number of RNAs per cell [14]. The final dataset therefore contained the normalized expression value, expressed as transcripts per cell, for 822 genes in 74 human cell types. Such a matrix is very useful in order to explore the expression profiles of various genes in different cell types. We therefore established these profiles for a number of genes (Figure 3). Two tendencies are immediately apparent from those patterns.

One is that the addition of null values (the absence of tags from some libraries) did not result in the generation of an aberrant point (Figure 3a-c). The pattern generated was very similar to the one generated for the tags belonging to the strictly minimal transcriptome set, that is, those for which no missing data were found (Figure 3d-e). All the patterns displayed a markedly skewed distribution toward the smallest values, which may directly result either from sequencing errors or from insufficient sequencing effort. This bias should be minimized when creating the boolean matrix (see below).

All the genes, whether belonging to the strictly minimal transcriptome set or not, display highly variable expression levels (ranging for example from 3 to 433 copies per cell for the GAPDH gene; Figure 3e). Such a high level of variation creates very interesting conditions for creating the boolean matrix. We verified that this very high dynamic range of GAPDH expression was not an artifactual result produced by large differences in library sizes by plotting the number of observed GAPDH tags as a function of library size (Figure 3g). Clearly, no correlation could be detected between these two parameters. Indeed, the very high variation in levels of GAPDH has recently been documented using both SAGE and quantitative RT-PCR [15].

The values contained in the 822 x 74 expression matrix had to be transformed into boolean values by a so-called discretization phase. In our context, each quantitative value has

given rise to one boolean value, that is, true (1) or false (0). We have explored the effect of three different discretization procedures (Figure 4). First we used the ‘max minus x%’ procedure. This consists of identifying the highest expression value (HV) in any library for each tag, and defining a value of 1 for the expression of that tag in a library when the expression value was above $(HV - x)/100$. Otherwise, the expression of the tag was assigned a value of 0 (Figure 4a).

We also analyzed the effect of a mid-range-based cutoff (Figure 4b). The highest and lowest expression values were identified for each tag and the mid-range value was defined as being equidistant from these two numbers (their arithmetic mean). Then, all expression values below or equal to the mid-range were set to 0, and all values strictly above the mid-range were set to 1.

Finally, we used a ‘x% of highest value’ approach (Figure 4c). For each tag, we identified libraries in which its level of expression was in the 5% of highest values. These were assigned the value 1, and the rest were set to 0.

These procedures will clearly generate different boolean matrices. One of the main questions will then be: what is the influence of those differences on the nature of the generated rules? As seen below, some tendencies are conserved irrespective of the discretization procedure applied; others are only apparent using one of the three approaches.

The use of such discretization approaches is nevertheless both technically interesting and biologically sound. The main technical interest lies in the elimination of all the very low tag values, which may be the result of either sequencing errors or insufficient sequencing effort. Biologically, the discretization approach can be seen as an overexpression experiment, in which we identify genes that are overexpressed (right part of the rule) when a given set of genes are overexpressed (left part of the rule). Nevertheless, differences that are generated as a result of different discretization approaches highlight the problem of determining the best mathematical definition of gene overexpression.

Mining the boolean context and post-processing the rules

The binary matrices were transformed into a readable format for the ac-miner software (see Materials and methods). For the extraction phase, in every case we applied a confidence value of 100%. This means that all the rules generated are true in 100% of the cases. We first checked the effect of varying the frequency threshold, from 10% to 2%, on the number of rules extracted (Table 2).

It is obvious from the large differences in the number at different frequency thresholds, that very different boolean values were generated in the various boolean matrices. The proper threshold giving a reasonable amount of rules has

Tag Sequence	Description
AAAAATTCT	Taa matches mitochondrial sequence
AAAAGAAACT	POLYADENYLATE-BINDING PROTEIN
AACGCCGCCA	Macrophage migration inhibitory factor
AACTAAAAAA	Ubiquitin A-52 residue ribosomal protein fusion product 1
AACTAACAAA	Ubiquitin A-52 residue ribosomal protein fusion product 1
AAGACAGTGG	Ribosomal protein L37a
AAGGAGATGG	Ribosomal protein L31
AAGGTGGAGG	60S RIBOSOMAL PROTEIN L18A
AATAGGTCCA	Ribosomal protein S25
AATCCTGTGG	Ribosomal protein L8
ACAGGGTGAC	Homo sapiens mRNA for EDF-1 protein
ACAGTGGGGA	Human (p23) mRNA, complete cds
ACATCATCGA	Ribosomal protein L12
ACTTTTTCAA	Taa matches mitochondrial sequence
AGAATCGCTT	Homo sapiens coatomer protein (COPA) mRNA, complete cds
AGCACCTCCA	Eukarvotic translation elongation factor 2
AGCCCTACAA	Taa matches mitochondrial sequence
AGCCCTCCCT	Homo sapiens autoantigen p542 mRNA, complete cds
AGCTCTCCCT	60S RIBOSOMAL PROTEIN L23
AGGAAAGCTG	ESTs. Highly similar to 60S RIBOSOMAL PROTEIN L36 [Rattus norvegicus]
AGGCTACGGA	60S RIBOSOMAL PROTEIN L13A
AGGGCTTCCA	UBIQUINOL-CYTOCHROME C REDUCTASE COMPLEX SUBUNIT VI REQUIRING PROTEIN
AGGTGGCAAG	Taa matches mitochondrial sequence
ATAATCTTTT	Ribosomal protein S29
ATCAAGGGTG	Ribosomal protein L9
ATCACGCCCT	Taa matches mitochondrial sequence
ATCAGTGGCT	PROTEASOME BETA CHAIN PRECURSOR
ATGAAACCCC	Homo sapiens mRNA expressed in osteoblast, complete cds
ATGGCTGGTA	40S RIBOSOMAL PROTEIN S2
ATTCTCCAGT	Ribosomal protein L17
ATTTGAGAAG	Taa matches mitochondrial sequence
CAAGCATCCC	Taa matches mitochondrial sequence
CAATAAACTG	PROTEIN TRANSLATION FACTOR SUI1 HOMOLOG
CAATAAATGT	Ribosomal protein L37
CACAAACGGT	40S RIBOSOMAL PROTEIN S27
CAGCAGAAGC	Homo sapiens 4F5rel mRNA, complete cds
CATCTAAACT	Human mRNA for KIAA0038 gene, partial cds
CCACCCCGAA	Testis enhanced gene transcript
CCACTCCTCA	DEFENDER AGAINST CELL DEATH 1
CCAGGAGGAA	HEAT SHOCK COGNATE 71 KD PROTEIN
CCAGTGGCCC	Ribosomal protein S9
CCATTGCACT	Ataxia telangiectasia mutated (includes complementation groups A, C and D)
CCCATCCGAA	Ribosomal protein L26
CCCAGCCAG	Ribosomal protein S3
CCCAGTTGC	Calpain, small polypeptide
CCCGTCCGGA	60S RIBOSOMAL PROTEIN L13
CCCTGGGTTT	Ferritin, light polypeptide
CCGGGTGATG	Human copper transport protein HAH1 (HAH1) mRNA, complete cds
CCGTCCAAGG	Ribosomal protein S16
CCGTGCTCAT	ESTs. Highly similar to ADIPOCYTE P27 PROTEIN [Mus musculus]
CCTAGCTGGA	PEPTIDYL-PROLYL CIS-TRANS ISOMERASE A
CCTCGAAAAA	60S RIBOSOMAL PROTEIN L38
CCTGGAAGAG	"Procollagen-proline, 2-oxodulurate 4-dioxygenase (proline 4-hydroxylase), beta polypeptide (protein disulfide isomerase)
CCTTGGAGAT	Ribosomal protein S5
CGCCGCGGGC	Human ribosomal protein L35 mRNA, complete cds
CGCCGCGGTG	Homo sapiens Chromosome 16 BAC clone CIT987SK-A-761H5
CGCCGGAACA	Ribosomal protein L4
CGCTGTTTCC	Homo sapiens ribosomal protein L11 mRNA, complete cds
CGTGTTAATG	CELLULAR NUCLEIC ACID BINDING PROTEIN
CTAAGACTTC	Taa matches mitochondrial sequence
CTAGCCTCAC	Actin, gamma 1
CTCATAAGGA	Taa matches mitochondrial sequence
CTGCTATACG	Ribosomal protein L5
CTGCTGAGTG	ESTs. Highly similar to HYPOTHETICAL 14.1 KD PROTEIN C31A2.02 IN CHROMOSOME I [Schizosaccharomyces pombe]
CTGGCTGCAA	Cytochrome c oxidase subunit Vb
CTGGGGTAAT	40S RIBOSOMAL PROTEIN S19
CTGTTGATTG	Heterogeneous nuclear ribonucleoprotein A1
CTGTTGGTGA	40S RIBOSOMAL PROTEIN S23
GAAAAATGGT	Laminin receptor (2H5 epitope)
GAACACATCC	Ribosomal protein L19
GAAGCAGGAC	COFILIN, NON-MUSCLE ISOFORM
GAAGTTATGA	T-COMPLEX PROTEIN 1, ALPHA SUBUNIT
GACGACACGA	Ribosomal protein S28
GACTCACTTT	Peptidylprolyl isomerase B (cyclophilin B)
GAGGGAGTTT	Ribosomal protein L27a
GCAAAACCCC	Homo sapiens tumor necrosis factor superfamily member LIGHT mRNA, complete cds
GCACAAGAAG	ESTs
GCAGCCATCC	Ribosomal protein L28
GCATAATAGG	Ribosomal protein L21
GCATAGGCTG	ELONGATION FACTOR TU, MITOCHONDRIAL PRECURSOR
GCCAGCCGAG	Human transcriptional corepressor hKAP1/TIF1B mRNA, complete cds
GCCCCTCCGG	ESTs. Weakly similar to TRANS-ACTING TRANSCRIPTIONAL PROTEIN ICPO [Bovine herpesvirus type 1 (strain k22)]
GCCGAGGAAG	Human mRNA for ribosomal protein S12
GCCGTGTCCG	Human ribosomal protein S6 mRNA, complete cds
GCCTCTCCCT	ESTs
GCCTGTGGG	Phospholipid hydroperoxide glutathione peroxidase
GCCTGTATGA	Ribosomal protein S24
GCCTTCCAAT	P68 PROTEIN
GCGAAACCCC	Human G protein-coupled receptor (STR122) mRNA, complete cds (signal)
GCGAAACCCCT	V-erb-b2 avian erythroblastic leukemia viral oncogene homolog 3 (alternative products) (signal)
GCTGGCTGGC	Homo sapiens chaperonin containing t-complex polypeptide 1, eta subunit (CctH) mRNA, complete cds
GCTTTATTTG	Human mRNA fragment encoding cytoplasmic actin, (isolated from cultured epidermal cells grown from human foreskin)

Figure 2 (see the next page for continuation of figure and the legend)

Tag Sequence	Description
AAAACATTCT	Taa matches mitochondrial sequence
GGACCACTGA	Ribosomal protein L3
GGAGTGGACA	60S RIBOSOMAL PROTEIN L18
GGCAAGAAGA	Ribosomal protein L27
GGCAAGCCCC	Heat shock 27kD protein 1
GGCCGGGTTT	Ribosomal protein S17
GGCCTGCTGC	ESTs. Highly similar to C10 [H.sapiens]
GGCTGGGGGC	Human profilin mRNA, complete cds
GGCTTTACCC	Eukarvotic translation initiation factor 5A
GGGCGCTGTG	Homo sapiens mRNA for smallest subunit of ubiquinol-cytochrome c reductase, complete cds
GGGGAATCG	THYMOSIN BETA-10
GGGGGACGGC	ESTs. Weakly similar to Y48E1B.1 [C.elegans]
GGGGGTACC	ATP SYNTHASE LIPID-BINDING PROTEIN P1 PRECURSOR
GGTCCAGTGT	Phospholipase mutase 1 (brain)
GGTGGATGTG	Homo sapiens methyl-CoG binding protein MBD3 (MBD3) mRNA, complete cds
GGTTTGGCTT	UBIQUINOL-CYTOCHROME C REDUCTASE COMPLEX 11 KD PROTEIN PRECURSOR
GTGAAACCTC	Homo sapiens intrinsic factor-B12 receptor precursor, mRNA, complete cds
GTGAAGGCAG	Ribosomal protein S3A
GTGACAACAC	Voltage-dependent anion channel 1
GTGACAGAAG	Eukarvotic translation initiation factor 4A (eIF-4A) isoform 1
GTGACCTCCT	CYTOCHROME C OXIDASE POLYPEPTIDE VIII-LIVER/HEART PRECURSOR
GTGATGGTGT	Thyroid autoantigen 70kD (Ku antigen)
GTGCTGAATG	MYOSIN LIGHT CHAIN ALKALI, SMOOTH-MUSCLE ISOFORM
GTGGCACACG	Eukarvotic translation initiation factor 3 (eIF-3) p36 subunit
GTGGCAGGTG	Human mRNA for KIAA0340 gene, partial cds
GTGGCTCACA	Adenosine A2b receptor
GTGTTAACCA	Human ribosomal protein L10 mRNA, complete cds
GTGTTGCACA	Ribosomal protein S13
GTTAACGTCC	Homo sapiens Bruton's tyrosine kinase (BTK), alpha-D-galactosidase A (GLA), L44L-like ribosomal protein (L44L) and FTP3
GTTGGTGCCA	Ribosomal protein L35a
GTTGTGGTTA	BETA-2-MICROGLOBULIN PRECURSOR
TAAGGAGCTG	Ribosomal protein S26
TAATAAAGGT	40S RIBOSOMAL PROTEIN S8
TACAAGAGGA	Ribosomal protein L6
TACCATCAAT	Glyceraldehyde-3-phosphate dehydrogenase
TACCTAGAA	Estrogen receptor
TAGGTTGTCT	TRANSLATIONALLY CONTROLLED TUMOR PROTEIN
TAGTTGAAGT	UBIQUINOL-CYTOCHROME C REDUCTASE COMPLEX 14 KD PROTEIN
TCACAAGCAA	H.sapiens alpha NAC mRNA
TCAGATCTTT	Ribosomal protein S4, X-linked
TCAGTTTGTCT	Human HS1 binding protein HAX-1 mRNA, nuclear gene encoding mitochondrial protein, complete cds
TCTCCAGGAA	ESTs. Weakly similar to PUTATIVE MITOCHONDRIAL CARRIER C16C10.1 [C.elegans]
TCTTGTCAT	L-LACTATE DEHYDROGENASE M CHAIN
TGAAGGAGCC	ATP SYNTHASE LIPID-BINDING PROTEIN P2 PRECURSOR
TGAGGGAATA	Triosephosphate isomerase 1
TGCACGTTTT	Human mRNA for antileukoprotease (ALP) from cervix uterus
TGCATCTGGT	78 KD GLUCOSE REGULATED PROTEIN PRECURSOR
TGGAATGCTG	Homo sapiens NADH:ubiquinone dehydrogenase 51 kDa subunit (NDUFB1) mRNA, nuclear gene encoding mitochondrial
TGGAGTGGAG	Human guanylate kinase (GUK1) mRNA, complete cds
TGGCTGTGTG	ESTs
TGGGCAAAGC	Translation elongation factor 1 gamma
TGGTGTGAG	Human DNA sequence from clone 1033B10 on chromosome 6p21.2-21.31. Contains the BING5 gene, exons 11 to 15 of the
TGTACCTGTA	Human alpha-tubulin mRNA, complete cds
TGTGATCAGA	Homo sapiens F1F0-type ATP synthase subunit alpha mRNA, complete cds
TGTGCTAAAT	60S RIBOSOMAL PROTEIN L34
TGTGTTGAGA	Translation elongation factor 1-alpha-1
TTATGGGATC	GUANINE NUCLEOTIDE-BINDING PROTEIN BETA SUBUNIT-LIKE PROTEIN 12.3
TTCATACACC	Taa matches mitochondrial sequence
TTCTTGTGGC	Ribosomal protein S11
TTGGAGATCT	Human NADH:ubiquinone oxidoreductase MLRQ subunit mRNA, complete cds
TTGGGGTTTC	Ferritin heavy chain
TTGGTCTCT	60S RIBOSOMAL PROTEIN L41
TTGTAATCGT	Human mRNA for ornithine decarboxylase antisense, ORF 1 and ORF 2
TTGTTGTTGA	Calmodulin 1 (phosphorylase kinase, delta)
TTTAACGGCC	Taa matches mitochondrial sequence

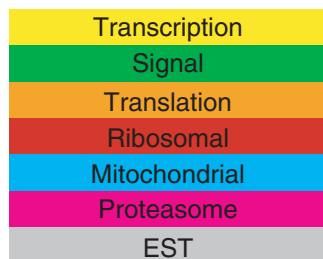


Figure 2 (continued from the previous page)

The strictly minimal transcriptome set, representing the 156 tags that were found in each and every SAGE library constructed from more than 20,000 sequenced tags. The key shows the color code used for representing the function of the encoded proteins.

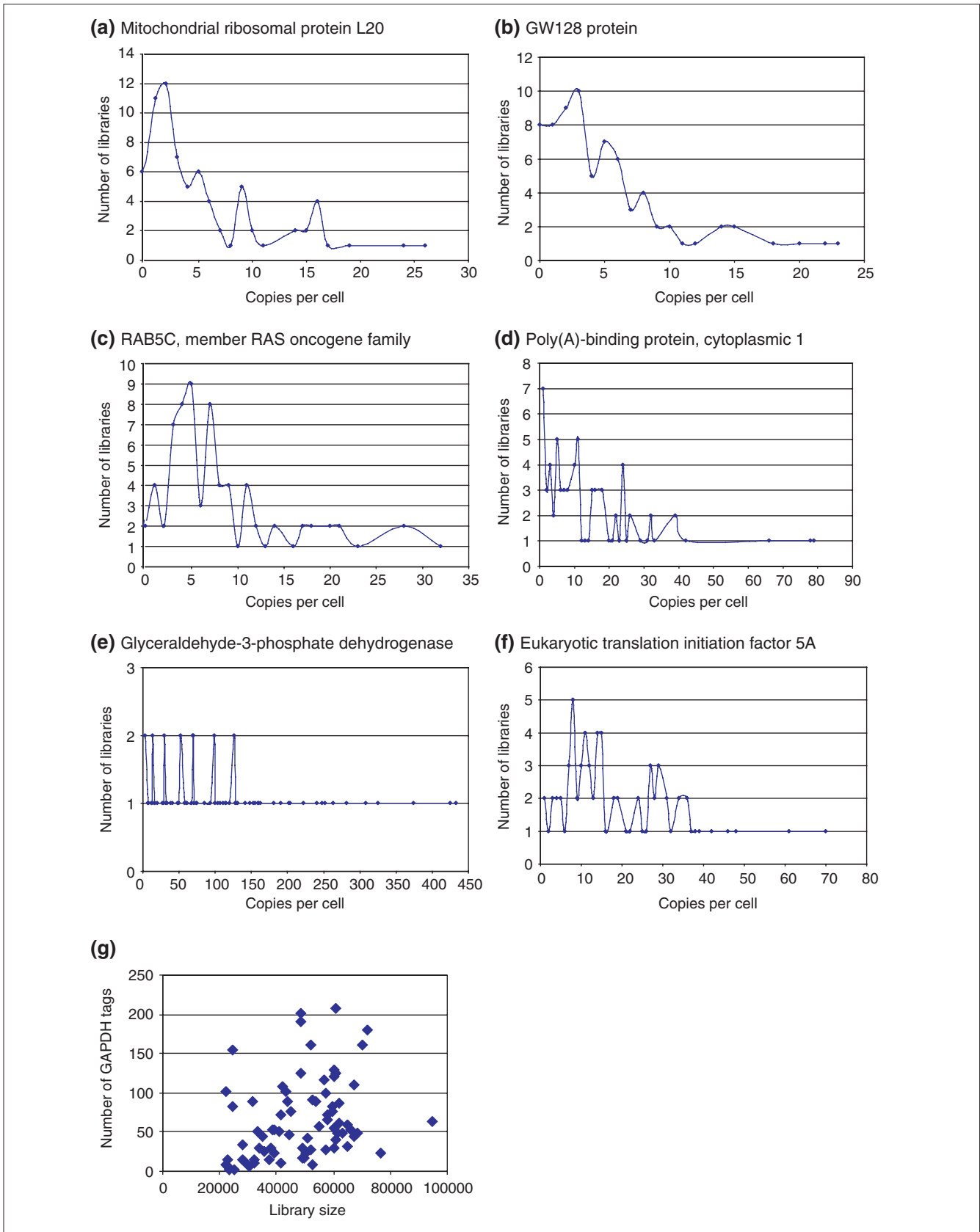
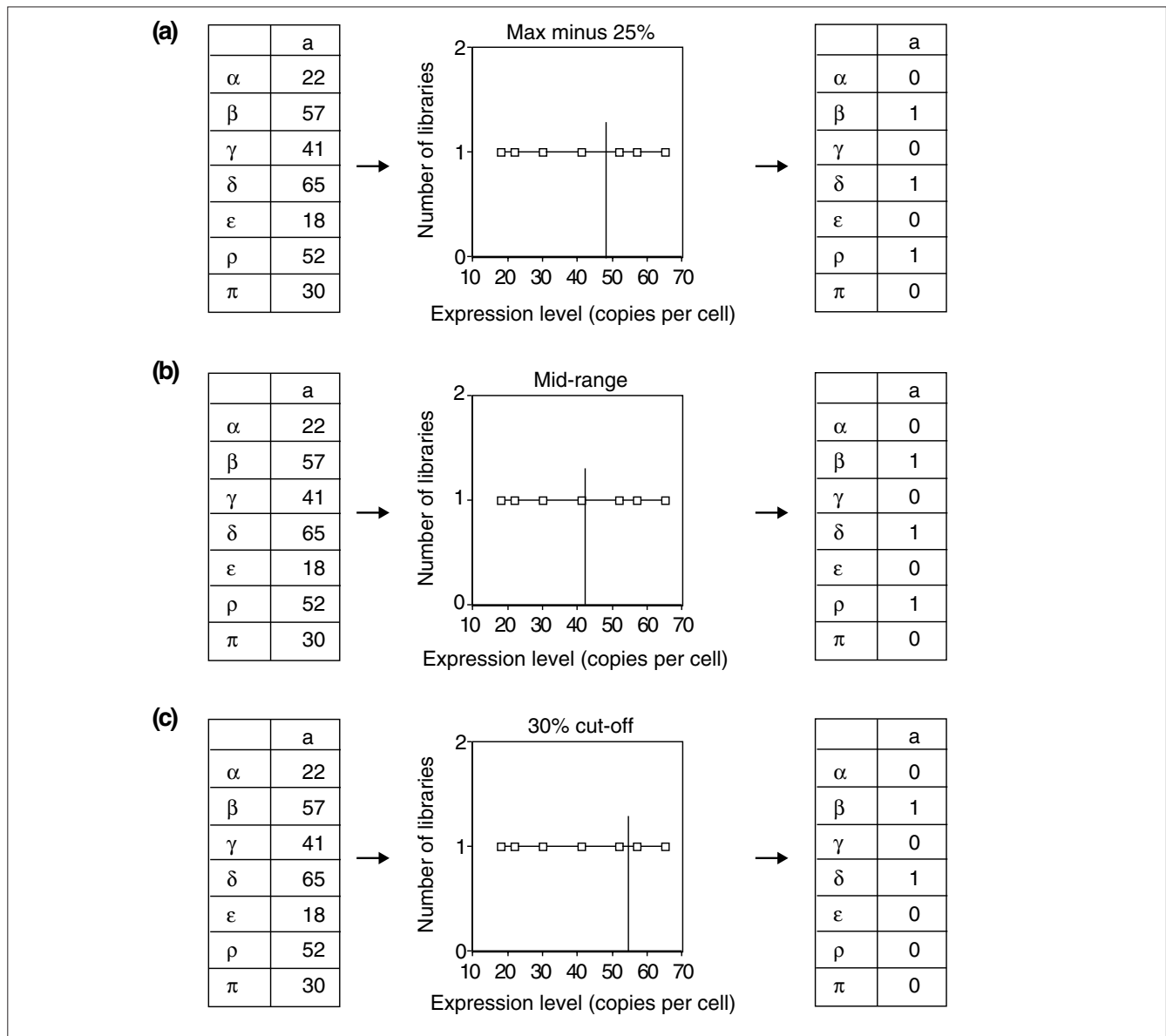


Figure 3 (see legend on the next page)

Figure 3 (see figure on the previous page)

Gene-expression profiles represented as copies per cell, derived by normalization from the values obtained in the 74 libraries used. **(a)** Mitochondrial ribosomal protein L20; **(b)** GWI28 protein; **(c)** RAB5C, member of the Ras family; **(d)** Poly(A)-binding protein, cytoplasmic I; **(e)** glyceraldehyde-3-phosphate dehydrogenase (GAPDH); **(f)** eukaryotic translation initiation factor 5A. **(g)** Repartition of GAPDH frequency (in number of tags) as a function of the size of the library from which that frequency was deduced. Three tags are shown that belong to the strictly minimal transcriptome set (d-f) and three tags not belonging to that set (a-c).

**Figure 4**

Schematic description of the discretization protocols used. On the left is shown the original matrix containing continuous values expressed as copies of transcript per cell, in the center the discretization procedure used, and on the right the resulting Boolean matrix. **(a)** The max minus 25% approach. We took the highest value (65) and calculated 25% of 65 (16.25). Values above $65 - 16.25 (= 48.75)$ were given a boolean value of 1; all others were given a value of 0. **(b)** The mid-range approach. We took the highest value (65), the lowest value (18), and calculated the mid-range as $((65-18)/2)+18= 41.5$. Values above 41.5 were given a boolean value of 1; all others were given a value of 0. **(c)** The 30% cut-off. We took the highest 30% of the number of values (here $7 \times 0.3 = 2.1$, that is, the two highest values), set them to 1 and the rest to 0. Note that this 30% value is not the actual value used for the extraction (see below) but is given as an illustration. Note also that these three different procedures resulted in different matrix densities. The last procedure resulted in the same number of boolean 1 results for all tags, whereas the first two generated different densities of 1, depending on the gene-expression pattern throughout the various libraries.

therefore to be determined empirically on a given boolean matrix. It can also be seen from this table that the number of rules generated for a low-frequency threshold can be so high that even a properly designed algorithm is unable to extract them. In some cases at a low threshold, although rules can be extracted, their very high number renders their analysis impossible, at least with the post-processing strategies and tools currently available.

The 68 rules extracted from the 'Max minus 25% matrix' using an ac-miner frequency threshold at 5% were then transferred into an Excel file and color-coded according to the function encoded by the genes associated in the rules (Figure 5). It was immediately obvious that we obtained a very high level of homogeneity in the rules, with most of the rules associating genes encoding ribosomal proteins.

We then analyzed the rules from the 'mid-range-based' cutoff using an ac-miner frequency threshold at 10%. The resulting 1,746 rules were transferred into an Excel file and color-coded according to the function encoded by the genes associated in the rules (see Additional data files). Once more, it was immediately obvious that we obtained a very high level of homogeneity in the rules, and once again most of the rules associated genes encoding ribosomal proteins. This time, a higher incidence of other translation-related non-ribosomal proteins was observed. It was also clear that because of the high number of rules generated, a post-processing task would need to be carried out to identify a smaller number of rules with the highest biological relevance. This is always the case with such unsupervised data mining techniques.

Finally, we analyzed the rules extracted from the '5% of highest value' matrix using two different ac-miner frequency

Table 2

Effect of varying the frequency threshold on the number of rules extracted

Matrix	Frequency threshold (%)	Number of rules
Max minus 25%	10	0
Max minus 25%	6	10
Max minus 25%	5	68 (Figure 5)
Max minus 25%	2	4,084
Mid-range-based	10	1,746
Mid-range-based	6	36,550
Mid-range-based	5	111,143
Mid-range-based	2	Intractable
5% of HV	10	0
5% of HV	6	21
5% of HV	5	301
5% of HV	2	32,329

thresholds. The 5% frequency value resulted in the generation of 301 rules that were transferred into an Excel file and color-coded according to the function encoded by the genes associated in the rules (see Additional data files). Once again, the rules were homogeneous, mostly associating protein synthesis-related (ribosomal or not) tags. Nevertheless, it was also apparent that some clustering of genes found in mitochondrial DNA was beginning to appear. This was made much more obvious by extracting rules at a 2% frequency value, and by selecting from the resulting 32,329 rules those that visually contained more than one tag identifying a mitochondrial gene. This approach resulted in a set of 436 rules that were highly homogeneous (see Additional data files).

Biological significance of the discovered rules

To explore the biological significance of the generated rules, we decided to investigate in detail a subset of the longest rules generated from the mid-range-based approach. We focused on a set of 47 rules associating 13 different genes (see Additional data files). As previously noted, the rules are very homogeneous, mostly associating protein synthesis-related tags. Only two tags were not categorized as such: a yellow-coded tag (number 763, transcription) and a gray-coded tag (631, expressed sequence tag (EST)). First, we rechecked the identity of tag 763 (sequence TGGTGTGAG), by using the 'tag to gene' function of the National Center for Biotechnology Information (NCBI) website [6]. It provided the identification: Hs.275865: ribosomal protein S18. The initial identification of this tag was thus misleading and this shows that our data-mining technique can allow the correct reassignment of wrongly labeled tags.

We then rechecked the identity of tag 631 (sequence GTTG-GTCTGT). It provided the following identification: Hs.288967: *Homo sapiens* cDNA FLJ14105 fis, clone MAMMA1001202. This is an EST of unknown function. The results obtained with our mining technique suggest that the putative protein encoded by this EST is involved in protein synthesis. The ability both to correctly relabel wrongly attributed tags and to propose a function for an unknown protein is a first element of biological validation of the generated set of rules, although final verification will depend on biological experiments demonstrating a function.

To gain further insight into the biological meaning of the rules generated, we decided to investigate in finer detail several rules associating proteins involved in signal transduction. The following rules were highlighted:

When the gene encoding the S6 kinase (RSK-B) (Tag n° 579) is overexpressed (that is, it has an expression value of 1), then the gene encoding the Lamin A (Tag n° 598) is also overexpressed, and this is true in 4 different cell types (see line 10 in Figure 5a for the actual the rule). This association of S6 kinase and lamin A is in good accordance with the previous identification of S6 kinase II as a potentially important lamin kinase [16].

When the gene encoding the ras-related GTP-binding protein (Tag n° 312), is overexpressed (that is, it has an expression value of 1), then the gene encoding the Platelet-endothelial tetraspan antigen 3 (Tag n° 730) is also overexpressed, and this is true in 8 different cell types. Tetraspans are membrane proteins that are proposed to act as 'molecular facilitators', grouping specific cell-surface proteins and thus increasing the formation and stability of functional signaling complexes [17]. Although they were proposed to activate the Jun N-terminal kinase (JNK) pathway [18], no association with Ras-family proteins has previously been reported. ARD results would support the contention that the function of Ras-family proteins is coordinated with that of tetraspan.

When both the genes encoding the G protein-coupled receptor (STRL22) (Tag n° 459) and the KH type splicing regulatory protein KSRP (Tag n° 795) are overexpressed, then the gene encoding the KIAA0340 gene (Tag n° 607) is also overexpressed, and this is true in 8 different cell types. This rule suggests the existence of a functional association between a G-protein-coupled receptor related to chemokine receptors [19], a splicing factor [20], and a member of the RAS gene superfamily (KIAA0340, also known as RAB3A, RIM, RIM1, RAB3IP2, RIMS1 [21]). That the expression of a G protein-coupled receptor correlates with that of a small GTP-binding protein may not be surprising. However, the indication that this signaling pathway may somehow be functionally correlated with a splicing event is an interesting prediction.

Comparing association rules with self-organizing maps

To compare the output of ac-miner with a traditional clustering method, we decided to use a recently described implementation of the SOM algorithm called SOTA, which is freely available on the web [22,23]. The expression matrix, before boolean transformation, was computed through SOTA, using a variability threshold of 90%, and the resulting clusters were color-coded (see Additional data files). As such, it is clear that this approach does not allow either relabeling of tags, or assignment of putative functions to tags encoding unknown proteins. Nevertheless, it can be quite useful to compare the results generated by the two methods, especially for exploring the rules associating heterogeneous members. One can for example note that one SOTA cluster associates several of the genes that were associated in the longest rules (Figure 6). The fact that four genes (including the EST of unknown function) belong at the same time to a strong rule and to a SOTA cluster clearly reinforces the prediction that those four genes are co-regulated and participate in a similar function.

Discussion

We have used a new data-mining technique (ARD) for analyzing gene-expression matrices to discover potentially interesting strong association rules. To validate the potential

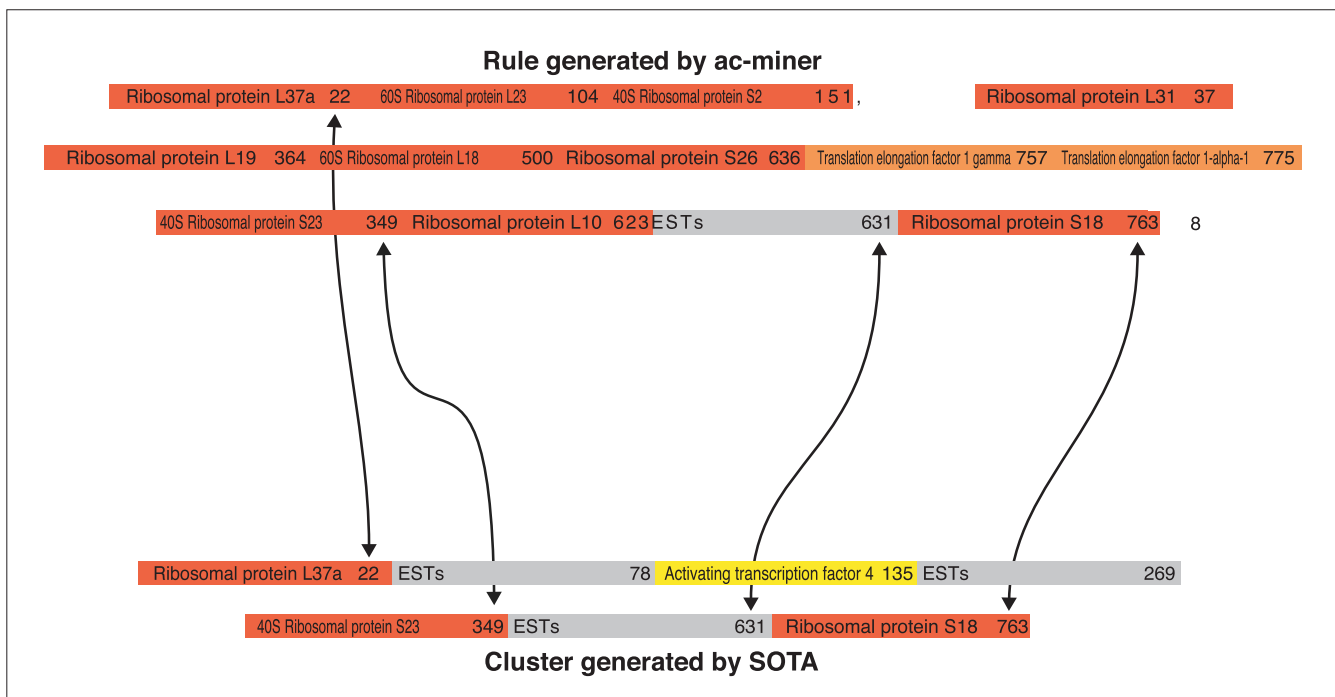
of this technique, we used expression data that are freely available (human SAGE data). Because of the very high dimensionality of SAGE data, application of the recently developed algorithm Min-Ex has proved very useful compared to the previously developed association-rule algorithms based on the Apriori algorithm [4].

This approach was successful in generating a set of rules that were biologically meaningful according to two main criteria. This first criterion was that the rules generated were very homogeneous in terms of the function associated with the tags. The very strong message revealed by our approach is that some ribosomal mRNAs are strongly co-regulated. A very similar finding has been recently described in various biological systems, including many tumors and cell lines [24-26], cells overexpressing erbB-2 [27], cells overexpressing N-myc [28], NGF-treated PC12 cells [29] and yeast cells examined at a compendium level [30]. Both the underlying mechanism(s) and the biological function(s) for such co-regulation are still essentially elusive (see discussion in [29]), especially in light of the described translational control of ribosomal protein synthesis [31]. It is nevertheless clear that our 'blind' mining technique essentially captured a biological phenomenon that until now has been described on dissimilar occasions, and could represent a more general phenomenon than anticipated. We also observed the expected co-regulation of genes encoded by mitochondrial DNA (see [32] for a recent review). We were also able to extract several rules involving signal transduction proteins, some of which were confirmed by previous published evidence, and some of which point in yet-unexplored directions.

The color-coding we used is somewhat 'crude' and finer biological definitions would be necessary for investigating the biological meaning of the rules in greater detail. We plan to use the newly described Gene Ontology classification [33] as a basis for establishing tag meta-identity. It would be very interesting to see whether rule homogeneity can be observed at the three GO levels: not only at the biological process level, but also at the molecular function and cellular component levels.

The homogeneity that was generated in the rules was sufficient to relabel a wrongly labeled tag, and to propose a putative function for an EST-encoded protein. Beyond the validation purpose, the heterogeneous sets of rules might be considered as the most biologically interesting, as they might point to previously unidentified functionally connected sets of genes. It is clear that the usefulness of such a set of rules for the scientific community will be very much improved by the use not only of the minimal transcriptome set, but of all the thousands of non-ambiguous tags (tags identifying only one gene and present at least once in two different libraries, see [9]). This work is in progress in our laboratories.

We are currently exploring other promising future directions. One is to evaluate a number of various objective mea-

**Figure 6**

Side-by-side comparison of one rule generated by ac-miner, belonging to the longest set and one cluster generated by SOTA (see Additional data files). The arrows indicate tags that were found to associate through both methods.

asures of interestingness to rank the rules according to criteria complementary to user-based knowledge. For example, conviction [34] is interesting, because, contrary to the confidence measure, it takes the support of the right-hand side of the rule into account. Roughly speaking, conviction measures the quality of a rule seen as an implication. The J-measure [35] measures two important criteria for the quality of a rule, namely simplicity and goodness-of-fit between the rule and the data. Interestingly, when applied to frequent rules, we need only know the frequent itemsets (see Materials and methods for a definition) and their frequencies. Thus these measures can be evaluated efficiently.

Another extremely interesting scientific challenge is to consider other user-defined constraints that can be inserted in the association-rule miner algorithms [36]. These would not only be related to frequency and confidence thresholds but also to various constraints, for example, those that enforce the presence or absence of some attribute. This might enable work at lower-frequency thresholds and a focus on rules that involve the desired attributes. This is important, as it might enable the identification of association rules that are not sufficiently strong to be discovered at present.

Another line of enquiry is the possibility of applying the association-rule approach to associate cell types rather than genes, in order to identify closely related cell types. Such an approach has already been implemented, using a clustering

algorithm on human SAGE data [9]. It remains to be seen whether the ARD approach can give rise to rules that produce cell regroupings that can be compared with these clusters, just as rules and clusters associating genes have been compared.

To conclude, we have demonstrated the usefulness of ARD, and in particular that this approach can be used to highlight various properties concealed in the mass of data, depending on the discretization procedure applied. This approach should be regarded as highly complementary to the more traditional clustering approaches, and should prove as useful for extracting properties from microarray-generated expression data as we demonstrated for SAGE-generated data.

Materials and methods

The association-rule discovery technique

The classical framework

In the standard presentation of ARD [4], the data are in the form of a (large) boolean matrix. In the case of gene-expression data, it may be generated from DNA arrays or SAGE approaches, and initial values are continuous. The generation of the boolean matrix from such continuous data is a critical step that may impinge on the interpretation of the rules generated. We have described three different approaches to generating such matrices (see 'Preprocessing raw data and preparing the boolean context') that all try to capture a gene-expression pattern.

If the columns of the matrix are denoted C_1, C_2, \dots, C_n , then the ARD technique discovers association rule that are expressions $X \Rightarrow Y$ where X and Y are sets of columns (often called itemsets) and $X \cap Y = \emptyset$ (for an example of the actual rules extracted, see Figure 5). The relevance of a rule $X \Rightarrow Y$ can be measured by its frequency and its confidence. The frequency is the number of rows where all columns in X and Y have a value of 1 (true) simultaneously. This number is often divided by the number of rows to provide a relative frequency. The confidence is the ratio between the frequency and the number of rows where all the columns in X have a value of 1, that is, it estimates the conditional probability of observing the properties denoted by Y when the properties denoted by X are true. Thus, when a rule has a confidence of 1, it means that 100% of the rows that have 1 in the columns in X also have 1 in the columns in Y . The classical association-rule mining task concerns the discovery of every rule such that its frequency and its confidence are greater or equal to user-defined thresholds [4].

Given our example dataset in Table 1 we define that gene a is overexpressed in condition β if its level of expression is greater than the mean level of expression of a across all conditions (α to π). Otherwise, it is considered to be underexpressed (see ‘Preprocessing raw data and preparing the boolean context’ above for the actual discretization procedures used). In this way we build a boolean matrix as shown in Table 3.

In the example shown above, the rule gene $a \Rightarrow$ gene c has the frequency $3/7$ and the confidence $3/4$. In other words, genes a and c are overexpressed simultaneously in $3/7$ of the situations, and when gene a is overexpressed then in 75% of the situations gene c is also overexpressed. If the frequency threshold had been fixed at 30% and the confidence threshold at 70%, then the rule gene $a \Rightarrow$ gene c would be discovered. At these thresholds, a rule such as gene $c \Rightarrow$ gene b would not be discovered because it is not sufficiently valid ($4/6 = 66\%$), although it is sufficiently frequent ($4/7 = 57\%$). Similarly, the rule gene $a, \text{ gene } b, \text{ gene } c \Rightarrow$ gene d would

not be discovered because it is not sufficiently frequent ($1/7 = 14\%$), although it is valid (confidence = 100%).

Mining of frequent and valid association rules has led to a great deal of research since the definition of the Apriori algorithm [4]. Many data-mining software tools include an implementation of this algorithm, for example, commercial packages such as Clementine [37] and free academic software packages such as Weka [38].

Solutions to the classical association-rule mining problem

Most of the available implementations of association-rule miners perform well on sparse and weakly correlated boolean data. They are based on a two-phase process. First, every frequent itemset is identified and its frequency computed. An example of an itemset would be the sets of genes that are overexpressed together in more than 10% of the biological situations. This will represent the limiting step, in terms of computing time, for the whole process.

Valid rules are identified from each discovered frequent itemset. The discovery of valid frequent rules from the frequent itemsets is a straightforward operation. For each frequent itemset X , the rule $X \setminus \{Y\} \Rightarrow Y$ is generated for every subset Y of X , and its confidence is calculated. Rules are listed only if their confidence is greater or equal to the user-defined threshold. Using this approach, the derivation of frequent/valid rules is quite efficient as infrequent itemsets have already been eliminated to reduce the numbers of calculations to determine validity and confidence of rules.

It turns out that the challenging issues related to association-rule mining are now being able to compute every frequent itemset and its frequency for the desired frequency threshold, and being able to support the identification of useful rules among the (thousands of) extracted frequent and valid rules. The identification of frequent itemsets in a boolean matrix consists of retrieving all possible attribute combinations that are true-valued in a number of lines of the matrix greater than the frequency threshold. For example in a matrix of 1,000 columns the initial number of candidates is $2^{1,000}$. The computation of frequent itemsets is only feasible if an efficient candidate-enumeration strategy enables the search space to be pruned of most of the infrequent subsets without having to count their frequency. Fortunately, one can use the so-called anti-monotonicity of the minimal frequency as a constraint. Thus, an itemset cannot be frequent if one of its subsets is not frequent. Thus by identifying first singletons, then pairs, then itemsets of size 3, and so on, it is possible to prune most of the search space. This is because as soon as an itemset is known to be infrequent, all its supersets can be ignored. By considering larger and larger itemsets, it is clear that eventually the whole collection of the frequent itemsets will be identified. In practice, apriori-based frequent-itemset-extraction algorithms work if the largest frequent itemset has a size lower or equal to 15 (at

Table 3

Boolean matrix built from the matrix of expression values				
	a	b	c	d
α	0	1	1	0
β	1	0	1	0
γ	1	1	0	0
δ	1	1	1	1
ϵ	0	1	1	1
ρ	1	0	1	0
π	0	1	1	1

least 2^{15} itemsets have been considered by the counting procedure). Clearly, the lower the frequency threshold, the larger the number of frequent itemsets.

A reasonable frequency threshold can be operationally defined as one at which the extraction of rules remains tractable (that is, does not require excessive computing time) but which nevertheless conserves sufficiently interesting properties. Thus the properties one wishes to study should have to involve perhaps at least 5% of the situations, whereas the extraction becomes intractable below 30%. We might want to compute the frequent sets for the lowest possible threshold to provide more input for the post-processing phase. Apriori-like implementations can fail because of this essential requirement to identify and count for at least every frequent itemset. In situations in which the number of frequent itemsets is high, the frequency threshold must be increased and the risk that the frequent patterns discovered are trivial also increases similarly.

The post-processing phase is important for reporting results and supporting the discovery of interesting rules among the frequent and valid rules. Taking into account the huge number of generated rules, we must provide not only tools allowing us to browse and select the rules (subjective criteria of interestingness) but also various objective measures of interestingness to rank the rules according to complementary criteria (see Discussion).

Furthermore, among the set of frequent and valid rules, a large number of rules will certainly be redundant. For example in the situation of rules $r_1 = X_1 \Rightarrow Y_1$ and $r_2 = X_2 \Rightarrow Y_2$ (when $X_1 \subset X_2$ and $Y_2 \subset Y_1$) are both frequent rules and have (almost) the same confidence. We can consider that r_1 is more general than r_2 and therefore discard r_2 from the output because r_2 does not provide any additional information. Application of this simple technique allows thousands and in some cases hundreds of thousands of redundant rules to be discarded without losing any information. Note, however, that this is an application-independent post-processing operation: no domain knowledge is used to eliminate uninteresting rules. In a complementary manner, domain knowledge can be applied to select subsets of generated rules, for example, the rules that contain some attributes could be discarded or attention can be focused on the subsets of rules in which others attributes occur.

With respect to the data described in Results, we have tried the classical approach and had difficulties using an Apriori-based implementation. Not only were we forced by constraints of tractability to work at rather high frequency thresholds, but despite this, in this exploratory context, we were nevertheless drowned by the number of rules generated. Instead of implementing tedious post-processing phases, we decided to apply a recent technique that proposes solutions to these two problems.

Mining rules in dense boolean matrices: the Min-Ex approach

A huge research effort has been made to solve these problems of Apriori-like algorithms. The proposals concern both the efficient computation of the frequent itemsets and the possibility of directly identifying rules of greater potential interest. The design of Min-Ex and its implementation *ac-miner* is the result of this international effort [39]. Not only does it enable the more efficient computation of frequent itemsets but it also directly identifies a subset of the frequent and valid rules in condensed representation of the frequent itemsets.

Min-Ex is based on the concept of δ -free itemsets. An itemset is designated δ -free if none of its subsets is linked together in a δ -strong rule, that is, a rule with at most δ exceptions. In the research reported here, we used only rules that were 100% true, that is $\delta = 0$. In this case, an itemset X is δ -free if there is no association rule with confidence of 100% between subsets of X . There are two very interesting properties of δ -free itemsets. First, anti-monotonicity of freeness; every subset of a δ -free itemset is also δ -free and every superset of a non δ -free itemsets is also non δ -free. Second, frequent δ -free itemsets are a condensed representation of frequent itemsets, that is, they are less numerous than the frequent itemset while providing almost the same information (exactly the same information if $\delta = 0$).

Taken together, those two properties allow that when X is known to be a frequent δ -free itemset, the frequency of (many) supersets of X can be derived from the frequency of X without having to count them. More precisely, for each δ -free set X , one can efficiently compute the maximal superset F such that its frequency is equal to the frequency of X , the so-called closure of X . As a result, every set that is included between X and F is known to have the frequency of X .

In real-life data, it enables one to reduce significantly the number of sets for which the counting phase is needed. Also, it reduces the size of the explored search space [39]. Formalization and technical algorithmic details for computing δ -free itemsets and their closures are described in [39]. In the present paper, we use the implementation called *ac-miner-close*.

Example

In the example boolean matrix, {gene a, gene b} is δ -free (there is no 100% confidence rule $\text{gene a} \Rightarrow \text{gene b}$ or $\text{gene b} \Rightarrow \text{gene a}$). {gene b, gene c, gene d} is not δ -free and its frequency can be derived from the frequency of {gene d} since there is a 100% confidence rule $\text{gene d} \Rightarrow \text{gene b, gene c}$.

The *ac-miner-close* association rule miner provides each frequent δ -free set, its frequency and its closure. For instance $\langle \text{gene d, gene b gene c: } 3 \rangle$ says that {gene d} is δ -free and is true in 3 lines (above the provided frequency threshold). Furthermore, it says that each time gene d is true (that is,

has a value of 1, and is thus considered as overexpressed), gene b and gene c are true as well. In other terms, we have a rule $\text{gene d} \Rightarrow \text{gene b} \text{ gene c}$ with a confidence value of 1 (logical rule). Thus, mining 0-free sets and their closures leads to association rules with confidence 1.

When considering the whole collection of logical rules extracted with ac-miner-close, we have an important property that these rules are not only frequent and logical (confidence 1) but maximal: $X \Rightarrow Y$ is maximal if another frequent and logical rule $Z \Rightarrow W$ such that $Z \subseteq X$ and/or $Y \subseteq W$ does not exist. Ac-miner-close provides the collection of such rules, designated strong rules, directly. These rules have a minimal left-hand side and a maximal right-hand side and many redundant rules are not provided.

Clearly, provided that the computation is tractable, which will be determined by the frequency threshold, the strong rules can also be computed from the output of an Apriori-based association rule miner. It needs for the computation of the so-called structural cover of the extracted rules (discarding rules that do not satisfy the above property of maximality).

Additional data files

Four supplemental figures are available with the online version of this paper. The rules in the table are color-coded according to the function encoded by the genes associated in the rules. For the color-coding conventions used in the supplemental figures, see Figure 2.

Supplemental figure 1 shows the first 566 rules (out of 1,746) generated by ac-miner on a mid-range-based boolean matrix using a 10% frequency threshold.

Supplemental figure 2 shows (a) The 301 rules generated by ac-miner on a '5% cut-off' boolean matrix using a 5% frequency threshold. (b) The 436 rules, out of 32,329 generated by ac-miner on a 5% cut-off boolean matrix using a 2% frequency threshold, and from which the rules containing mitochondrial tags were visually extracted.

Supplemental figure 3 shows the longer set of rules: from the 1,746 rules generated by ac-miner on a mid-range-based boolean matrix using a 10% frequency threshold. Those 49 rules associated 13 tags and were true in 8 libraries. Shown are the rules (a) before and (b) after the proper reassignment of the tag 763.

Supplemental figure 4 shows the SOTA clustering output translated into a color-coded Excel file.

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- SAGE data: description files** [<ftp://ftp.ncbi.nlm.nih.gov/pub/sage/info/>]
- SAGE data: Stats.txt** [<ftp://ftp.ncbi.nlm.nih.gov/pub/sage/OLD/extra/stats.txt>]
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