THE USE AND ANALYSIS OF MICROARRAY DATA

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Functional genomics is the study of gene function through the parallel expression measurements of genomes, most commonly using the technologies of microarrays and serial analysis of gene expression. Microarray usage in drug discovery is expanding, and its applications include basic research and target discovery, biomarker determination, pharmacology, toxicogenomics, target selectivity, development of prognostic tests and disease-subclass determination. This article reviews the different ways to analyse large sets of microarray data, including the questions that can be asked and the challenges in interpreting the measurements.

At the time of writing this article, the genomes of more than 800 organisms have been sequenced, and well over 3.5 million genetic sequences have been deposited in international repositories. However, the biological functions of most of these genes remain unknown, or have been predicted only through homology to genes with functions that are better known. One way to determine the functions of these genes is through repeated measurements of their RNA transcripts; for example, knowing that a particular gene is expressed only in cardiac muscle and only under particular conditions implicitly gives us functional knowledge about that gene. Functional genomics is the study of gene function through parallel expression measurements of a genome. The most common tools used to carry out these measurements include complementary DNA microarrays¹, oligonucleotide microarrays² or serial analysis of gene expression (SAGE)3. This article will focus on microarrays, which are artificially constructed grids of DNA, such that each element of the grid probes for a specific RNA sequence — that is, each holds a DNA sequence that is a reverse complement to the target RNA sequence.

Although there are many protocols and types of system available, the basic technique involves extraction of RNA from biological samples in either normal or interventional states. The RNA (or, in some protocols, isolated messenger RNA) is then copied, while incorporating either fluorescent nucleotides or a tag that is later stained with fluorescence. The labelled RNA is then

hybridized to a microarray for a period of time, after which the excess is washed off and the microarray is scanned under laser light. This process is schematized in FIG. 1. With oligonucleotide microarrays, for which all probes have been designed to be theoretically similar with regard to hybridization temperature and binding affinity, each microarray measures a single sample and provides an absolute measurement level for each RNA molecule, although this absolute measurement might not correlate exactly with concentration in terms of micrograms per unit volume. With cDNA microarrays, for which each probe has its own hybridization characteristic, each microarray measures two samples, and provides a relative measurement level for each RNA molecule. Regardless of the technique, the end result is 4,000-50,000 measurements of gene expression per biological sample. As a complete experiment might involve anywhere up to hundreds of microarrays, the resultant RNA-expression data sets can vary greatly in size.

As the cost of microarrays continues to drop, it is clear that microarrays are becoming more integral to the drug discovery process. In addition to the obvious use of functional genomics in basic research and target discovery, such as finding genes expressed in significantly different patterns across samples, there are many other specific uses in this domain. These include: biomarker determination, to find genes that correlate with and presage disease progression, but are easier to measure and follow in clinical trials; pharmacology, to

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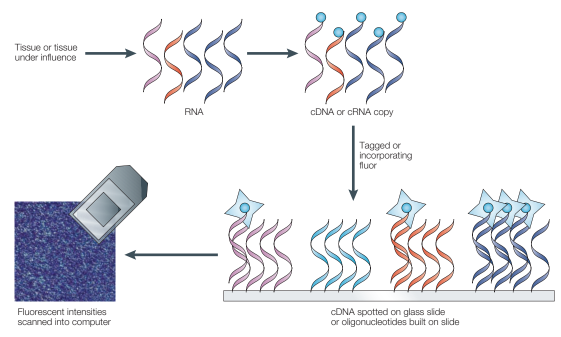


Figure 1 | **Schematized experimental process using a microarray**. Although the specific protocols differ, the microarray approach first involves isolating RNA or messenger RNA from appropriate biological samples, making the RNA (or a copy of it) fluorescent, hybridizing it to the microarray, washing off the excess and scanning the microarray under laser light. cDNA, complementary DNA; cRNA, complementary RNA.

determine differences in gene expression in tissues exposed to various doses of compounds; toxicogenomics, to find gene-expression patterns in a model tissue or organism exposed to a compound and their use as early predictors of adverse events in humans; target selectivity, to define a compound by the gene-expression pattern it provokes in a target tissue and then compare it with other compounds using these patterns; prognostic tests, to find a set of genes that accurately distinguishes one disease from another; and disease-subclass determination, to find multiple subcategories of tumours in a single clinical diagnosis.

Many free (BOX 1) and commercial software packages are now available to analyse microarray data sets, although it is still difficult to find a single off-the-shelf software package that answers all functional-genomics questions. As the field is still young, when developing a bioinformatics analysis pipeline, it is more important to have a good understanding of both the biology involved and the analytical techniques rather than having the right software. This article reviews the different ways to analyse microarray data, and will concentrate on choosing the appropriate method for the given hypothesis.

Normalization and noise

Before multiple microarray measurements can be integrated into a single analysis, the reported measurements need to be normalized, or modified (possibly corrected) to make them comparable. When microarrays are used to collect gene-expression data in an experiment in which the measurements are made at the same time, with homogeneous populations of similar cells and using a

single microarray technology, normalization might simply be a matter of adjusting the overall brightness of each scanned microarray image, assuming that the quantity of RNA is equal⁴. Other normalization methods include: using expression levels of 'housekeeping' genes⁵; using assumptions that most genes do not change across experiments⁶; using SPLINES⁷; or other nonlinear techniques^{8,9}.

Typically, however, functional-genomics experiments are more complicated. Recently, increasing efforts have been invested in characterizing the 'noise' in microarray technology. Studies addressing the reproducibility of microarray data analysed replicated data¹⁰, compared microarray measurements with NORTHERN BLOTS^{11,12} and SAGE¹³, and evaluated strategies for REVERSE TRANSCRIPTION¹⁴ and *in vitro* transcription amplification¹⁵. As a result, it has become increasingly clear that there are several substantial sources of noise in microarray data. Intra- and inter-microarray variations can markedly skew the interpretation of such expression data.

First, improving the reliability of expression measurements starts with proper experimental design. For example, microarrays can measure across the genome, including genes with expression that is controlled by hormones, such as growth hormone or cortisol. So, if organ samples are acquired at various times during the day, genes that appear to be differentially expressed might only be reflecting normal circadian physiology. Pooling samples before hybridization might control for this biological 'noise.'

In addition, scanned hybridization images need to be inspected for artefacts, such as scratches and bubbles ^{16,17}. Measuring replicate microarrays for each biological sample allows the modelling of this technical noise.

SPLINES
Instead of fitting a complex
polynomial curve to data, splines
allow the fitting of data by
putting together smaller, less
complex curves.

NORTHERN BLOT
Different RNA molecules are
separated by mass on a gel, then
radioactively labelled
complementary DNA or RNA
molecules are used to quantify
specific RNA amounts.

REVERSE TRANSCRIPTION
The synthesis of a strand of
DNA from RNA, which is used
to make a complementary DNA
copy of sample RNA.

Most reported expression data have been obtained on relatively homogeneous cell populations. However, when RNA is extracted from whole organs or from tumour biopsies, the sources of variation increase. There is substantial heterogeneity of expression in cell subpopulations in most organs and in many tumours. Failure to account for such variation could lead to overinterpretation or spurious functional gene association. Microdissection of cell subpopulations (for example, with laser capture18) is possible only in a minority of the systems of interest. If microarray-based geneexpression measurements are to be reliable and economical, both at the level of basic biology and clinical assays, then all of these further sources of noise/variation must be incorporated directly into the analytical tools that interpret these data.

A further issue that needs to be addressed is the difference between the two most commonly used microarray technologies: spotted cDNA microarrays, which report differences in gene expression between two samples, and oligonucleotide microarrays, which report absolute expression levels. Normalization techniques for one microarray technology might not apply to another, owing to differences in assumptions and the distributions of the output measurements. For example, if we assume that in any given experiment, most genes in a cell do not change in expression and an equal number of genes are up- and downregulated (not always a valid assumption), then differential-expression measurements from spotted arrays might be found to be normally distributed, whereas measurements from oligonucleotide microarrays will not have the same distribution.

Expression measurements made across microarray technologies are not directly comparable. For example, the published microarray measurements from the National Cancer Institute (NCI) 60-cancer-cell-line panel, from spotted arrays¹⁹ and from oligonucleotide

arrays²⁰, show poor correlation between measurements²¹. This can be explained by the differences in low-level hybridization and analysis between the two techniques. To be precise, Affymetrix microarrays contain between 11 and 20 pairs of oligonucleotide probes for a target RNA, for which one of the pair is the reverse complement to an ideally unique 25-mer in the RNA and the other contains a mutated middle base pair and serves as a measure of stray signal. Using the differences between these intensities, the Affymetrix quantitative software judges the reliability of each probe pair and calculates a qualitative and quantitative measurement (see Affymetrix web site). Other quantitative methods are available in addition to the Affymetrix software⁷.

Compared with this, cDNA microarrays contain a single probe for each target RNA, and the two biological samples are different colours, so that after hybridization, the two colours are scanned separately and relative expression is determined by comparing intensities. There is striking non-correlation between Affymetrix quantitative measurements and ratios of intensities from a cDNA microarray, because these two technologies clearly measure expression differently.

Supervised or unsupervised

Current methodologies to analyse RNA-expression data sets can be divided into two categories: supervised approaches, or analysis to determine genes that fit a predetermined pattern; and unsupervised approaches, or analysis to characterize the components of a data set without the a priori input or knowledge of a training signal.

Supervised methods. Supervised methods are generally used for two purposes: finding genes with expression levels that are significantly different between groups

$\operatorname{Box} 1$ | Some freely available software for microarray analysis

Although many bioinformatics companies sell software that assists in microarray analysis, there are several freely available software packages that can be used to perform the six analytical techniques described in this article. Only a few are listed here.

 Although it is the standard for hierarchical clustering and viewing dendrograms, this software also creates selforganizing maps and performs principal-components analysis.

GeneCluster 2.0http://www-genome.wi.mit.edu/cancer/software/genecluster2/gc2.html

• Initially used for constructing self-organizing maps, the latest version now also finds nearest neighbours and performs other supervised methods. Written in Java, this program can essentially run under any computer operating system

• Software that creates self-organizing maps and performs hierarchical clustering, as well as finding principal components. This package also includes a component for support vector machines, but at present offers little for documentation. The software is written in Java, and a license for the source code of the software is also available.

MAExplorer. http://maexplorer.sourceforge.net/

This tool performs many aspects of microarray processing, including the raw image analysis. It contains a few
analytical techniques, including hierarchical clustering. The software is written in Java, and the source code is freely
available for modification.

*Software to create relevance networks. The software is written in Java, and a license for the source code is also available.

of samples, and finding genes that accurately predict a characteristic of the sample. Most functional-genomics experiments still typically use only a handful of microarrays (or equivalent technology), with samples measured under two or three conditions, and the application has the goal of finding those genes that are significantly differentially expressed. Significance has been evaluated in many different ways, including parametric22 and nonparametric tests^{23,24}, analysis of variance²⁵ and many others. Although it would be an understatement to call the analyses of these smaller sets of microarray data trivial, there are several published techniques that have been used to find genes that have the most different expression levels between a few samples. When determining whether a particular gene is differentially expressed between two samples, there are four characteristics that need to be considered: absolute expression level, or whether the gene is expressed at a high or low level; subtractive degree of change between groups, or the difference in expression levels across samples (calculated using subtraction); fold change between groups, or the ratio of expression levels across samples (calculated by division); and reproducibility of the measurement, or whether samples with similar characteristics have similar amounts of the gene transcript.

These four characteristics are related; for example, genes measured at low amounts of expression often have measurements that are less reliable, which leads to poor reproducibility across samples and high-fold changes that do not adequately describe the actual degree of change. All of the available techniques for comparing two sets of microarray measurements essentially evaluate these four characteristics for each gene in various ways to rank genes that are most significantly different.

For larger data sets, comparing microarrays one pair at a time misses trends that might exist between measurements. There are several published supervised methods that find genes or sets of genes that accurately predict sample characteristics, such as distinguishing one type of cancer from another, or a metastatic tumour from a non-metastatic one. These methods might find individual genes, such as the nearestneighbour approach²⁶, and/or multiple genes, such as decision trees²⁷, neural networks²⁸ and support vector machines^{29–31}. This article will focus on the two more popular supervised techniques: nearest-neighbour analysis and support vector machines.

Unsupervised methods. Users of unsupervised methods try to find internal structure or relationships in a data set instead of trying to determine how best to predict a 'correct answer'. Within unsupervised learning, there are three classes of technique: feature determination, or determining genes with interesting properties without specifically looking for a particular a priori pattern, such as principal-components analysis^{32–36}; cluster determination, or determining groups of genes or samples with similar patterns of gene expression, such as nearestneighbour clustering^{26,37}, self-organizing maps^{38,39}, *k*-means clustering and one- and two-dimensional hierarchical clustering^{19,40}; and network determination,

or determining graphs representing gene–gene or gene–phenotype interactions using Boolean networks^{41–43}, BAYESIAN NETWORKS⁴⁴ and relevance networks^{20,45,46}. This article will focus on the four most common unsupervised techniques of principal-components analysis, hierarchical clustering, self-organizing maps and relevance networks.

Like, but how different?

It is crucial to make a distinction between dissimilarity measures (also known as 'metrics') and clustering methods. A dissimilarity measure indicates the degree of similarity between two genes. A clustering method builds on these dissimilarity measures to create groups of features with similar patterns.

A commonly used dissimilarity measure is Euclidean distance, for which each gene is treated as a point in multidimensional space, each axis is a separate biological sample and the coordinate on each axis is the amount of gene expression in that sample^{35,38}. A schematic of this method is shown in FIG. 2a. One disadvantage of Euclidean distance is that if measurements are not normalized, correlation of measurements can be missed, the focus being instead on the overall amount of expression. A second disadvantage is that genes that are negatively associated with each other will be missed. Negative associations include gene interactions, such as those of tumour-suppressor genes. As an example, the tumourrepressor protein p53 acts as a transrepressor of several other genes. This means that the higher the level of p53, the less expression of other genes is expected. The concept of negative interaction is clearly different than the concept of no interaction.

Another dissimilarity measure that is commonly used is the Pearson Correlation Coefficient, which is measured between two genes that are treated as vectors of measurements⁴⁰ (FIG. 2b). The disadvantages in using this measure with gene-expression measurements are: first, it assumes that the measurements are normally distributed, which might not be the case for oligonucleotide-microarray measurements; and second, it assumes that genes interact in the assumed linear model, when in biology, a particular gene might best regulate other genes when in the middle of its own range of expression. Operationally, this measure is sensitive to outliers, and although techniques such as the RANK CORRELATION COEFFICIENT deal with these by replacing the measurements with ranks, it is not clear whether eliminating the outliers is ideal — many past discoveries have been found by focusing on the outliers in biology.

A third dissimilarity measure is mutual information (FIG. 2c), which allows for any possible model of interaction between genes and uses each expression-level measurement equally regardless of the actual value, and is therefore not biased by outliers⁴⁶. However, calculating the mutual information requires using discrete expression measurements (for example, representing the gene as 'high' and 'low', or 'high', 'medium' and 'low', and so on), and the mutual information depends on the number of 'bins' used. Ideally, this would be performed in a gene-specific manner, but sufficient information about

BAYESIAN NETWORK
A graphical representation
in which variables (that is,
genes) are represented as nodes.
Arrows between nodes represent
conditional dependence, which
is interpretable as causal
associations.

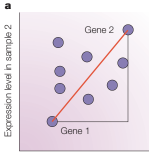
COEFFICIENT

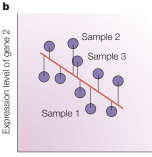
A measurement of the degree of fit of a linear-regression line to data points, calculated as the average distance of points from

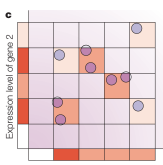
PEARSON CORRELATION

to data points, calculated as the average distance of points from the regression line normalized to the standard deviations of the individual coordinates.

RANK CORRELATION
COEFFICIENT
Points are restated in terms of
their ordinal rank (for example,
first, second, third) before
calculation of the correlation







Expression level in sample 1

Expression level of gene 1

Expression level of gene 1

Figure 2 | **Dissimilarity measures.** Almost all clustering techniques require some method of comparing one gene with another and determining similarity. **a** | Euclidean distance can be used as a measurement of the degree of similarity between two genes, and is calculated using the Pythagorean theorem. **b** | The Pearson correlation coefficient is calculated from the distances of each point from the linear-regression line (known as residuals). **c** | Mutual information is a model-free measurement of the degree of information content in one gene known from another gene, and is highest when genes are randomly distributed separately, but show a non-random joint distribution.

the range of expression of each gene in all tissues has not yet been collected. Furthermore, gene—gene associations with high mutual information might not even be functions in the mathematical sense, and might be difficult to explain biologically.

Analytical methods

Once a dissimilarity measure has been chosen, the appropriate analytical technique can be applied. This section describes the four commonly used unsupervised techniques — hierarchical clustering, self-organizing maps, relevance networks and principal-components analysis — and two commonly used supervised techniques — nearest neighbours and support vector machines.

Hierarchical clustering. Hierarchical clustering is a commonly used unsupervised technique that builds clusters of genes with similar patterns of expression. This is done by iteratively grouping together genes that are highly correlated in terms of their expression measurements, then continuing the process on the groups themselves. DENDROGRAMS (FIG. 3a) are used to visualize the resultant hierarchical clustering. A dendrogram represents all genes as leaves of a large, branching tree. Each branch of the tree links two genes, two branches or one of each. Although construction of the tree is initiated by connecting genes that are most similar to each other, genes added later are connected to the branches that they most resemble. Although each branch links two elements, the overall shape of the tree can sometimes be asymmetric. In visually interpreting dendrograms, it is important to pay attention to the length of the branches. Branches connecting genes or other branches that are similar are drawn with shorter branch lengths. Longer branches represent increasing dissimilarity.

Hierarchical clustering is particularly advantageous in visualizing overall similarities in expression patterns observed in an experiment, and because of this, the technique has been used in many publications^{40,47}. The number and size of expression patterns within a data set can be estimated quickly, although the division of the tree into actual clusters is often performed visually.

It is important to note the few disadvantages in their use. First, hierarchical clustering ignores negative associations, even when the underlying dissimilarity measure supports them. Negative correlations might be crucial in a particular experiment, as described above, and might be missed. Furthermore, hierarchical clustering does not result in clusters that are globally optimal, in that early incorrect choices in linking genes with a branch are not later reversible as the rest of the tree is constructed. So, this method falls into a category known as 'greedy algorithms,' which provide good answers, but for which finding the most globally optimal set of clusters is computationally intractable. Despite these disadvantages, hierarchical clustering is a popular technique in surveying microarray expression patterns in an experiment.

Self-organizing maps. Self-organizing maps are similar to hierarchical clustering, in that they also provide a survey of expression patterns within a data set, but the approach is quite different^{38,39}. As shown in FIG. 3b, genes are first represented as points in multidimensional space. In other words, each biological sample is considered a separate dimension or axis of this space, and after the axes are defined, genes are plotted using expression levels as coordinates. This is easiest to visualize with three or less microarrays, but extends to a larger number of experiments/dimensions. Nearness can be defined using any of the dissimilarity measures described above, although Euclidean distance is most commonly used.

The process starts with the answer, in that the number of clusters is actually set as an input parameter. A map is set with the centres of each cluster-to-be (known as centroids) arranged in an initial arbitrary configuration, such as a grid. As the method iterates, the centroids move towards randomly chosen genes at a decreasing rate. The method continues until there is no further significant movement of these centroids.

The advantages of self-organizing maps include easy two-dimensional visualization of expression patterns and reduced computational requirements compared with methods that require comprehensive pairwise

DENDROGRAM A visual representation of hierarchical clusters.

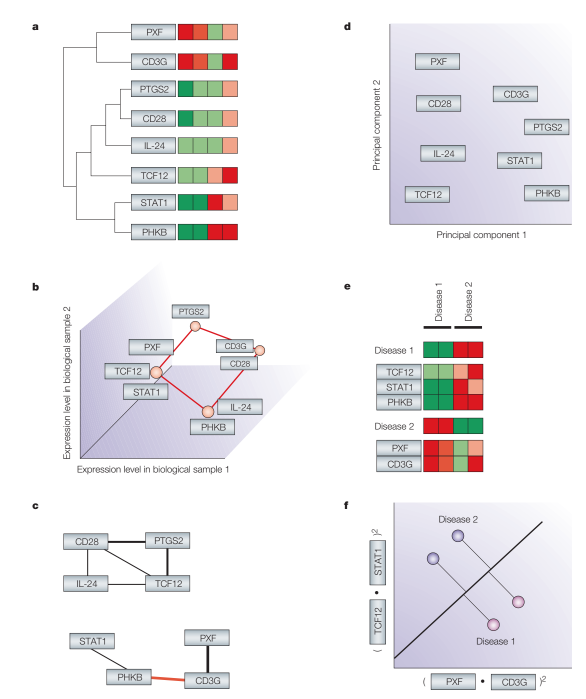


Figure 3 | Clustering and network-determination methods used in microarray analysis. The choice of the proper method and the results obtained clearly depend on the starting hypothesis. This figure shows the results of six analytical methods applied to the same hypothetical data set a | Hierarchical clustering sorts all genes (or samples), such that similar genes appear near each other. The length of the branch is inversely proportional to the degree of similarity. Shades of red indicate increased relative expression; shades of green indicate decreased relative expression. b | Self-organizing maps find variable-sized clusters of genes that are similar to each other, given the input number of clusters to find. c | Relevance networks find and display pairs of genes with strong positive and negative correlations, then construct networks from these gene pairs; typically, the strength of correlation is proportional to the thickness of the lines between genes, and red indicates a negative correlation. d | Principal-components analysis is typically used as a visualization technique, showing the clustering or scatter of genes (or samples) when viewed along two or three principal components. In the figure, a principal component can be thought of as a 'meta-biological sample', which combines all the biological samples so as to capture the most variation in gene expression. e | The nearest-neighbour supervised method first involves the construction of hypothetical genes that best fit the desired patterns (for example, a gene with high expression in disease 1 and low expression in disease 2, or vice versa). The technique then finds individual genes that are most similar to the hypothetical genes. f | Instead of restricting to individual genes, support vector machines efficiently try several mathematical combinations of genes to find the line (or plane) that best separates groups of biological samples. CD3G, CD3G antigen, γ-polypeptide; CD28, CD28 antigen; IL-24, interleukin-24; PHKB, phosphorylase kinase-8; PTGS2, prostaglandin-endoperoxidase synthase 2; PXF, peroxisomal farnesylated protein; TCF12, transcription factor 12; STAT1, signal transducer and activator of transcription 1.

comparisons, such as dendrograms. However, there are several disadvantages. First, the initial topology of a self-organizing map is arbitrary and the movement of the centroids is random, so the final configuration of centroids might not be reproducible. Second, similar to dendrograms, negative associations are not easily found. Third, even after the centroids reach the centres of each cluster, further techniques are needed to delineate the boundaries of each cluster. Finally, genes can belong to only a single cluster at a time.

Relevance networks. Continuing through the set of unsupervised techniques, relevance networks allow networks of features to be built, whether they represent genes, phenotypic or clinical measurements²⁰. The technique works by first comparing all features with each other in a pairwise manner, similar to the initial steps of hierarchical clustering. Typically, two genes are compared with each other by plotting all the samples on a scatterplot, using expression levels of the two genes as coordinates. A correlation coefficient is then calculated, although any dissimilarity measure can be used. A threshold value is then chosen, and only those pairs of features with a measure greater than the threshold are kept. These are displayed in a graph similar to FIG. 3c, in which genes and phenotypic measurements are nodes, and associations are edges between nodes. Although the threshold is chosen using permutation analysis, it can actually be used as a dial, increasing and decreasing the number of connections shown.

There are several advantages in using relevance networks. First, they allow features of more than one data type to be represented together; for example, if strong enough, a link between systolic blood pressure and expression of a particular gene could be visualized. Second, features can have a variable number of associations; theoretically, a transcription factor might be associated with more genes than a downstream component. Finally, negative associations can be visualized as well as positive ones.

One disadvantage to this method is the degree of complexity seen at lower thresholds, at which many links are found associating many genes in a single network. Completely connected subcomponents of these complex graphs (known as 'cliques') are not easy to find computationally.

Principal-components analysis. Principal-components analysis is more useful as a visualization technique than as an analytical method^{33,36}. It can be applied to either genes or samples, which are represented as points in multidimensional space, similar to self-organizing maps. Principal components are a set of vectors in this space that decreasingly capture the variation seen in the points. The first principal component captures more variation than the second, and so on. The first two or three principal components are used to visualize the gene on screen or on a page, as shown in FIG. 3d.

Because each principal component exists in the same multidimensional space, they are linear combinations of the genes. For example, the greatest variation of

biological samples might be described as 3 times the expression level of the first gene, plus –2.1 times the expression level of the second gene, and so on. The principal components are linear combinations that include every gene or sample, and the biological significance of these combinations is not directly intuitive.

There are other caveats in using principal components. First, it is important to note whether genes have been centred before analysis; if not, then the first principal component might serve to centre the genes. Second, it is crucial to note that although principal components might best describe the variation seen in an expression data set, they do not describe how to best separate groups of genes or samples. For example, if microarrays are measured on samples from two conditions, principal components will best describe the variation of those samples, but will not always be the best way to split samples from those two conditions⁴⁸.

Nearest neighbours. Although the nearest-neighbour technique can be used in an unsupervised manner, it is commonly used in a supervised fashion to find genes directly with patterns that best match a designated query pattern. For example, an ideal gene pattern might be one that is highly expressed in one condition and expressed at a low level in another condition. All the genes that have been measured can then be compared to this ideal gene pattern and ranked by their similarity, as shown in FIG. 3e. For example, acute lymphocytic leukaemia was distinguished from acute myelogenous leukaemia using this method²⁶.

Although this technique results in genes that might individually split two sets of microarrays, it does not necessarily find the smallest set of genes that most accurately splits the two sets. In other words, a combination of the expression levels of two genes might split two conditions perfectly, but these two genes might not necessarily be the top two genes that are most similar to the idealized pattern.

This technique can be modified for specific cases, such as toxicogenomics. Tissue exposed to various compounds known to cause different types of toxicity can be subjected to microarray measurements, as well as measurements from normal tissues. This could make up a training set, in that these data points implicitly make up a model of toxicity. Newer compounds can then be tested on these tissues (the test set), and the 'distance' of these expression patterns from the training set can be calculated, and a decision made as to similarity of mechanism of toxicity.

Support vector machines. Support vector machines address the problem of finding combinations of genes that better split sets of biological samples³⁰. Although it is easy to find individual genes that split two sets with reasonable accuracy owing to the large number of genes (also known as features) measured on microarrays, occasionally it is impossible to split sets perfectly using individual genes. The support vector machines technique actually further expands the number of features available by combining genes using mathematical

Box 2 | Downloadable large data sets of microarray measurements

Although Perou and others have called for the release of raw microarray data after the publication of manuscripts, this still rarely occurs⁵². The following web sites are the exception, and contain large amounts of microarray data that are of good quality and are freely available for academic use:

Stanford Microarray Database. http://genome-www5.stanford.edu/MicroArray/SMD/
• 3,290 Microarrays measured across 11 species, from 80 publications.

National Center for Biotechnology Information Gene Expression Omnibus...... http://www.ncbi.nlm.nih.gov/geo/
• 2,354 Microarrays from 105 types of microarray, measured across 78 experiments.

 ${\bf \cdot 565\,Microarrays\,from\,mouse\,and\,rat\,models\,of\,sleep, in fection, hypertension\,and\,pulmonary\,disease.}$

• 142 Microarrays involving mouse models of cardiac development and signal transduction, including measurements made in time-series.

• 121 Microarrays from 19 normal human tissues.

 $White head\ Institute\ Center\ for\ Genome\ Research\ http://www-genome.wi.mit.edu/cgi-bin/cancer/datasets.cgi-bin/c$

• Microarrays from 12 publications involving many types of cancer, including some clinical measurements associated with each sample.

operations (called kernel functions). For example, in addition to using the expression levels of two individual genes A and B to separate two sets of biological samples, the combination features $A \times B$, A/B, $(A \times B)^2$ and others, can also be generated and used. To make this clear, it is possible that even if genes A and B individually could not be used to separate the two sets of biological samples, together with the proper kernel function, they might successfully separate the two.

This can be visualized graphically as well, as shown in FIG. 3f. Similar to the principal-components analysis method above, consider each biological sample as a point in multidimensional space, in which each dimension is a gene and the coordinate of each point is the expression level of that gene in the sample. Using support vector machines, this high-dimensional space gains even more dimensions, representing the mathematical combinations of genes. The goal for support vector machines is to find a plane in this high-dimensional space that perfectly splits two or more sets of biological samples. Using this technique, the resulting plane has the largest possible margin from samples in the two conditions, therefore avoiding data over-fitting.

It is clear that within this high-dimensional space, it is easier to separate samples from two or more conditions, but one problem is that the separating plane is defined as a function using all the dimensions available. For example, the most accurate plane to separate one disease from another might be $(A \times B)^2 < 20$, where A and B are expression levels of genes. Although this might not be the most mathematically accurate way to separate two diseases, the biological significance of such functions is not always intuitive.

Relating hypotheses to techniques

As microarrays now cover the measurable genome, a user does not need to have a set of candidate genes in mind before use. However, this does not mean that a microarray user operates in a hypothesis-free manner. With all the supervised and unsupervised methods that are available for analysis, the challenge is in translating hypotheses into an appropriate bioinformatics technique.

In the domains of drug discovery and diagnostic testing, it is easy to find uses for supervised methods. Two examples illustrate this point. First, hypotheses in toxicogenomics, such as 'some genes in the genome influence liver metabolism of a particular compound', could be answered with a technique designed to find those genes for which expression measurements are most significantly different between liver cells with and without the compound. If many samples are present from each group, a nearest-neighbour approach could be used, in which an idealized gene is created that is expressed at a high level in samples with the compound and at a low level in samples without the compound. This is then used as a query to find genes that are most similar to the idealized pattern.

Second, hypotheses in the development of diagnostic tests, such as 'a combination of gene expression measurements accurately distinguishes metastatic from non-metastatic disease', can be answered by searching for such patterns using any of the supervised methods, including support vector machines. Certainly, any specific question can be answered directly in a supervised manner.

The need for unsupervised methods is less intuitive, because these start with a less direct question. For example, questions about the number and type of

expression responses in a period of time after application of a compound cannot be found in a supervised manner. Unsupervised techniques, such as hierarchical clustering and self-organizing maps, survey all genes and cluster them together on the basis of their expression patterns. A search for the pairs of genes that are most likely to be co-expressed can be accomplished using a technique such as relevance networks. Finally, true genetic regulatory networks (that is, hypotheses that the expression of one gene correlates with the expression of another) might be found using methods such as constructing Bayesian networks. What is common in these three examples is that there is no ideal answer that is being sought, and that it is difficult to ascertain when the method is correct. Nonetheless, unsupervised methods could be instrumental in the early discovery process.

There are a few special cases worth noting. When attempting to find connections between dissimilar items, such as a response to a small molecule (that is, a phenotypic measurement) and expression measurements, two-dimensional hierarchical clustering can be used for each measurement separately (for example, clustering the small molecules separately from the genes, as done by Ross *et al.*¹⁹), or building a relevance network (as in Butte *et al.*²⁰). Finally, when searching for subtypes of a condition or disease that might influence survival or a disease-free state, unsupervised hierarchical clustering can be paired with Kaplan–Meier survival statistics, as shown by Alizadeh and colleagues⁴⁹.

Challenges after analysis

After several microarray analyses, it quickly becomes obvious that the rate-limiting step in functional-genomics experiments is neither the handling of the biological samples nor the actual analysis, but instead the post-analytical work in determining what the results actually mean. First, detailed names and information might not yet be available for genes that have been found to be significant, even though these genes might have been measured on microarrays for years. This complicates the interpretation of results. The official gene name, predicted protein domains or gene-ontology⁵⁰ classification might become available as early as tomorrow, or as late as decades from now.

There are further post-analysis challenges. Occasionally, microarray probes are designed against chromosomal regions instead of expressed products, and when these probe sets are positive in an analysis, it is usually not clear which genes are being detected. It is worth finding these probe sets before analysis begins and eliminating them. Occasionally, probe sets

are incorrectly designed against the wrong strand or wrong species. Oligonucleotide sequences that were once thought to be unique for a particular gene might not remain unique as more genomic data are collected. Finally, in using spotted cDNA arrays, particularly those for which the probe sequences have not been validated, the findings might be incorrect.

Operationally, this means that one is never done analysing a set of microarray data. The infrastructure has to be developed to re-investigate constantly genes and gene information from microarray analyses performed in the past. It could be next month, for example, that new information about a gene that was positive in an analysis performed three months ago finally leads to a new and important hypothesis.

Conclusion

The challenge in determining the proper analytical methods to use is usually only a short-term difficulty, and typically, after the 'functional-genomics pipeline' has been established, the rate-limiting step shifts to the post-analytical challenges⁵¹. In the future, truly showing a 'return on investment' from functional genomics will depend on taking findings beyond the microarray stage and integrating them with the rest of the discovery pipeline. The 'list of genes' resulting from a microarray analysis should not be viewed as an end in itself; its real value increases only as that list moves through biological validation, ranging from the numerical verification of expression levels with alternative techniques, to ascertaining the meaning of the results, such as finding common promoter regions or biological relationships between the genes. However, tools that link these genes back to known biological pathways, as well as discovering new pathways, are in their infancy. Tools that can automatically indicate the importance of particular findings have yet to be invented. Until they come into being, the analysis of microarray data sets in a vacuum devoid of biological knowledge will be less rewarding.

Finally, the use of microarrays in basic and applied research in drug discovery is only going to increase, but as these data sets grow in size, it is important to recognize that untapped information and potential discoveries might still be present in existing data sets (BOX 2). It should be clear that any set of microarray measurements could be analysed and re-analysed in many different ways. In the application of functional genomics to drug discovery, to extract the most information from microarrays, an open mind always needs to be kept with regard to the choices of analytical methods, using supervised and unsupervised techniques, and methods yet to come.

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