# INTERACTIVE EXPLORATION OF MICROARRAY GENE EXPRESSION PATTERNS IN A REDUCED DIMENSIONAL SPACE

Jatin Misra<sup>1</sup>, William Schmitt<sup>1</sup>, Daehee Hwang<sup>1</sup>, Li-Li Hsiao<sup>2</sup>, Steve Gullans<sup>2</sup>, George Stephanopoulos<sup>1</sup>, and Gregory Stephanopoulos<sup>1\*</sup>

Department of Chemical Engineering (1) Massachusetts Institute of Technology Cambridge, MA 01890

And

Renal Division (2)
Department of Medicine
Brigham and Women's Hospital
Harvard Medical School
77 Avenue Louis Pasteur
Boston, MA 02115

### \*Corresponding author:

Gregory Stephanopoulos Department of Chemical Engineering, Room 56-469 Massachusetts Institute of Technology Cambridge, MA 02139

Phone: 617-253-4583, Fax: 617-253-3122

e-mail: gregstep@mit.edu

March 19, 2002

#### **ABSTRACT**

The very high dimensional space of gene expression measurements obtained by DNA microarrays impedes the detection of underlying patterns in gene expression data and the identification of discriminatory genes. In this paper we demonstrate the use of projection methods such as Principal Components Analysis (PCA) to obtain a direct link between patterns in the genes and patterns in samples. This feature is useful in the initial interactive pattern exploration of gene expression data and data-driven learning of the nature and types of samples. Using oligonucleotide microarray measurements of 40 samples from different normal human tissues, we show that distinct patterns are obtained when the genes are projected on a twodimensional plane spanned by the loadings of the two major principal components. These patterns define the particular genes associated with a sample class (i.e., tissue). When used separately from the other genes, these class- (i.e., tissue-) specific genes in turn define distinct tissue patterns in the projection space spanned by the scores of the two major principal components. In this study, PCA projection facilitated discriminatory gene selection for different tissues and identified tissue-specific gene expression signatures for liver, skeletal muscle, and brain samples. Furthermore, it allowed the classification of nine new samples belonging to these three types using the linear combination of the expression levels of the tissue-specific genes determined from the first set of samples. The application of the technique to other published data sets is also discussed.

**Keywords:** Projection methods, pattern discovery, gene expression, DNA microarrays, principal component analysis

#### INTRODUCTION

DNA microarrays are presently used extensively for genome-wide gene expression measurements. Large-scale transcriptional studies have catalyzed new discoveries and are generating important new insights into the behavior and functioning of cells (Alizadeh, *et al.*, 2000, Perou, *et al.*, 1999, Spellman *et al.*, 1998, Hughes *et al.*, 2000). Class discovery tools have played a key role in this process. Class discovery methods are exploratory analysis tools used to organize, learn from, and discover patterns in the data. Of the various multivariable techniques available, clustering of genes and samples has been the most common tool used for the analysis of microarray data (Alizadeh, *et al.*, 2000, Perou, *et al.*, 1999, Spellman *et al.*, 1998, Hughes *et al.*, 2000, Eisen *et al.*, 1998, Tamayo *et al.*, 1999). Before proceeding to cluster, it is often advantageous to visualize the data in order to develop an understanding of underlying structure. This initial exploration is useful in revealing patterns, and providing clues for further analysis.

Principal Component Analysis (PCA) is a linear projection method that defines a new dimensional space that captures the maximum information present in the initial data set by minimizing the error between the original data set and the reduced dimensional data set. Each principal direction of the projection space, or principal component (PC) is defined such as to be orthonormal to all others and to maximize the information in the data that has not already been captured by the previous (lower) dimensions. In this way, as the number of PCs progressively increases, a larger fraction of the total information content is accounted for. PCA is a linear projection in the sense that the variables of the projection space (PCs) are linear combinations of the original variables (i.e., the gene expressions). The coefficients of this linear combination are termed *loadings* and the

actual values of the projection of the samples are termed *scores*. PCA is obtained from a singular value decomposition of the data, and the loadings are the entries in the singular vector and are associated with genes. The scores are contained in the matrix obtained from a multiplication of the original data matrix with the singular vectors, and are associated with samples. Standard formulas are available for the determination of the projection variables, loadings and captured variability (Dillon and Goldstein, 1984), and many applications of PCA have been reported in a variety of different contexts (Alter *et al.*, 2000, Holter *et al.*, 2000, Rannar *et al.*, 1998, Kamimura, 1997).

In this paper we use PCA to analyze a set of microarray measurements on normal human tissues. Initial projection onto a lower dimensional space allows for better visualization of the entire data set. The loadings are subsequently used to select relevant genes, while considering the impact of the removal of irrelevant genes on the patterns observed in the projection of the samples. This is an alternate approach to the problem of selection of relevant genes in the analysis of microarray data (Golub *et al.*, 1999), and may be used to obtain a subset of genes that best describe the data. The observation of clear gene-expression patterns, after the removal of irrelevant genes, points to a high degree of structure in the measurements. Exploration of these gene expression patterns further revealed tissue-specific gene expression signatures. These signatures were further supported by the analysis of additional tissue samples that had not been used in the initial pattern discovery step.

#### RESULTS

The data set used in this study comprised expression measurements of 7070 genes made in 40 normal human tissue samples using Affymetrix GeneChips. The data were generated at the Brigham and Women's Hospital (BWH) in Boston (Hsiao *et al.*, 2001). Samples from several human tissues were analyzed, here we use the samples from brain, kidney, liver, lung, esophagus, skeletal muscle, breast, stomach, colon, blood, spleen, prostate, testes, vulva, proliferative endometrium, myometrium, placenta, cervix and ovarv.

#### PCA loadings can be used to filter irrelevant genes

The data from the 40 human tissues were first projected using PCA. PCA may be used with or without scaling (mean-centering, or autoscaling, among others). Here, we did not scale the data, and comparisons with mean-centered results are provided in the Discussion. The first and second PCs, account for approximately 70% of the information present in the entire data set. The score plot of the 40 samples using the entire gene expression set is shown in Fig. 1a. Also plotted in Fig. 1b are the loadings for each of the 7070 genes for the first and second PCs. The loading plot reveals a large number of genes clustered around the origin, implying that they only marginally impact the projection onto the first and second PC.

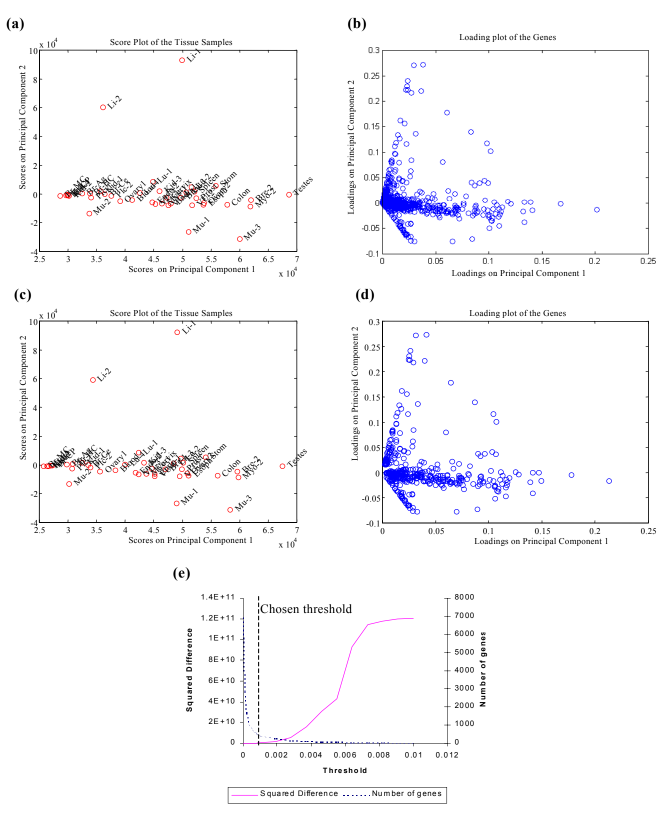


Figure 1: Gene selection based on the loadings on the principal components. Graphs (a) and (b) show the score plot of the samples and the loading plot of the genes, respectively, before any filtering is implemented. Graphs (c) and (d) show the score and loading plots after the filtering. Graph (e) displays quantitatively the decisions that went into the choice of the filtering threshold. It displays the distortion in the observed patterns, as measured through the squared difference, and the number of genes retained for analysis as the threshold is varied. The chosen filter threshold was 0.001. Filtering reduces the number of genes from 7070 to 425. At the same time, the score plot of the samples remains largely unchanged, and displays the same initial patterns, signifying a minimal loss of information. The loading plot displays strong linear structures of genes.

Since the relative magnitude of the loading is a measure of the importance of the corresponding gene in defining the PC, a small magnitude implies that the corresponding gene expression does not materially impact that particular PC. On this basis, a filter that eliminates genes with loadings below a threshold in all of the first five PCs was implemented. The decisions that went into the choice of the threshold are illustrated in Figure 1e. The threshold was varied over a large range, and at each threshold value, a record was maintained of the number of genes retained for analysis and the distortions in the score plot due to the elimination of genes. As the threshold value was gradually increased, the samples were re-projected using the subset of genes passing the filter. The distortion from the original score plot was measured in terms of the squared difference, defined as the sum of the squares of the difference between the 40 original score values and the 40 score values produced with the filtered gene set (this is defined mathematically in the Methods). In essence, this squared difference measures the error between the original projections and the new sample projections (or the distortion of the original pattern) as more and more genes are removed. When the threshold value exceeded 0.001, a large fraction of the genes were filtered out, precipitating large distortions in the patterns on the score plot. This criterion eliminated all but 425 genes whose loadings in at least one of the first five PCs exceeded the threshold value. A projection of the samples using only these 425 genes reveals an almost identical pattern on the score plot with the one obtained when all 7070 genes were used (Fig. 1c). This suggests that the dramatic reduction from the initial 7070 genes to the 425 finally retained resulted in a minimal information loss relevant to the description of the samples in the reduced space. Thus, a PCA framework may be used to evaluate the effect of gene removal on expression patterns observed in the reduced dimensional space.

## Identification of tissue-specific gene expression patterns. Correspondence between score and loading plots.

Three linear structures can be identified in the loading plot of the 425 genes selected by the above analysis, each structure comprising a set of genes arranged along a particular angle in Fig. 1d. These linear structures suggest a certain degree of organization in gene expression reflected in the linear relationships between the loadings of the first and second PCs of the genes clustered in these structures. An obvious question is whether there is any correlation among the genes that define these structures. Figure 2 shows the results of a systematic exploration of the patterns depicted in Fig. 1d. Plotted in Fig. 2a are the angles defined by the x-axis and the points representing the loadings of the first two PC's for the 425 consequential genes identified above. This histogram defines three clusters each corresponding to the three structures identified in Fig. 1d. The first, termed structure A, comprises genes with angles between 1.452 - 1.469 radians. The second, structure B, is centered around the second peak with angles between -1.222 and -1.205 radians, and the third is a set of genes between -0.328 and 0.054 radians, called structure C. The list of genes so selected was further refined to prevent the inclusion of genes that may have the same angle but are far removed from the structures in Fig. 1d by clustering the genes on the basis of their distance from the origin (the clustering results are discussed and provided in the Supplementary Materials). The final list of selected genes is provided in Table 1.

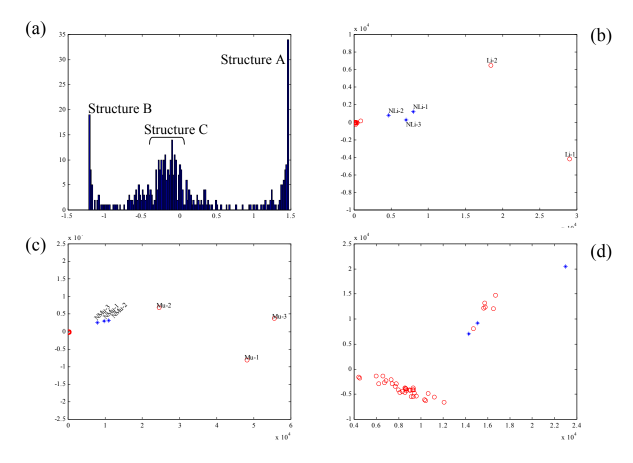


Figure 2: Identification of tissue-specific genes and validation using new samples. (a) Histogram of the angles between the x-axis and the points defined by the two principal loadings of each gene shown in Fig. 1d. Three main features, corresponding to the linear structures shown in Fig. 1d can be discerned, and are labeled as A, B and C. (b) PCA projection of all samples using the genes in Structure A. The samples in the initial data set are represented by red circles, and the new samples by blue asterixes. The two liver samples in the initial data set (Li-1, Li-2) and the new liver samples (NLi-1, NLi-2, NLi-3) are separated from the other samples, all of which cluster at the origin. (c) Projection of all samples using the genes in structure B. The muscle samples in the initial data set (Mu-1, Mu-2, Mu-3) are separated from the other samples along PC1. All the other tissue samples cluster at the origin. The new muscle samples are also separated when projected using these genes (NMu-1, NMu-2, NMu-3). (d) Projection of all samples using the genes in structure C. The six brain samples in the initial data set, and the three new brain samples are separated from the other samples.

Although the identity of some genes in the above groups are suggestive of the type of tissue they represent (for example, the genes in structure A contain an excess of genes related to the liver, such as albumins and apolipoproteins), the nature of each gene group is revealed when score plots are constructed using *only* the genes that are specific to the structures of Fig. 1d or 2a. Thus, using only the 24 genes of structure A to project all the samples yields a score plot (Fig. 2b) that dramatically separates the two liver samples in the data set from all the remaining tissue samples. Similarly, projecting the expression data of the 19 genes in structure B separates the three skeletal muscle tissue samples from the remaining tissues along the first PC (Fig. 2c) and, finally, projection of the samples using the 86 genes of structure C separates all six brain samples from the remaining tissues (Fig. 2d).

Inspection of the genes in structure C revealed two broad classes of genes. One class of genes with low expression levels was largely related to ribosomal proteins and function, while the other class of genes with larger and more variable expression are primarily brain tissue related genes. The loadings of these genes on the second PC support this observation, so that genes with high expression levels in the brain samples also had a high loading magnitude on the second PC, as shown in Table 1. This is also true of the genes in the other structures. This fact may be used for class discovery and data-driven learning, and is a result of the observed correspondence between the score plot and the loading plot. Given the observed separation of the six brain samples on the second PC in Fig. 2d, a learning approach for samples with unidentified characteristics would have consisted of the following steps: select a set of genes with high loadings on the dominant PC, examine their function, and generate hypotheses as to the nature of the samples. This is a class-discovery approach, in contrast to a classification methodology that relies on a priori labeling of the samples (Golub, et al., 1999, Brown et al., 2000). Here, the methodology allows one to probe the nature of the sample, and simultaneously identify the genes that contribute to the differentiation of the sample(s) from the others.

The genes that were not part of these structures were also analyzed, by projecting the samples using these genes, however, no clustering of samples or any noteworthy separation was observed.

<u>Table 1:</u> List of genes identified by angle selection. The genes are sorted by their loadings on the PC which separates the specific tissue. Also provided is the ratio of the mean of the gene expression in the specific tissue sample to the mean of the gene expression in all the other tissues. Genes with large values of the ratio tend to have large PC loadings. In the case of the brain-specific signature, only the top 30 genes as ranked by their loads on PC 2 are provided. A complete list of genes is in Supplementary Materials.

Gene ID	Ratio of Means	Loading	Gene Description
Liver Specific Signature	Tyreans	PC 1	
M36803	213.5	0.3293	hemopexin
J02843	337.8	0.3284	cytochrome P450IIE1 (ethanol-inducible)
X53595	344.5	0.318	beta-2-glycoprot I (apolipoprot H)
HG2841-HT2970	197.3	0.3175	Albumin 5
HG2841-HT2969	161.5	0.3042	Albumin, 3
M13149	131.5	0.2592	histidine-rich glycoprot
M10050	291.6	0.2533	liver fatty acid binding prot (FABP)
X03168	2313.7	0.2242	S-prot
D14446	148.2	0.2113	HFREP-1
M16961	161.2	0.2067	alpha-2-HS-glycoprot alpha and beta chain
X51441	342.2	0.1958	serum amyloid A (SAA) prot, clone pAS3-alpha
HG1827-HT1856	284.2	0.1956	Cytochrome P450, Subfamily Iic
L00190	254.4	0.1614	D29832, M21642 and others
M58600	1225.6	0.1523	heparin cofactor II (HCF2)
M21642	183.9	0.1265	(dysfunctional) antithrombin III (ATIII) Utah
M19828	1577.6	0.1064	apolipoprot B-100 (apoB)
M11567	3034.8	0.1059	angiogenin and three Alu repetitive sequences
X14690	222.4	0.1045	plasma inter-alpha-trypsin inhibitor heavy chain H(3)
M21642	128.9	0.096	(dysfunctional) antithrombin III (ATIII) Utah

M20786	248.8	0.0929	aluha 2 ulagmin inhihitan
M11321	317.2	0.0929	alpha-2-plasmin inhibitor group-specific component vitamin D-binding prot
U08006		0.0881	
J03474	146.8		complement 8 alpha subunit (C8A)
	132.6	0.0778	transcription factor SP1
S48983	358.8	0.0771	SAA4=serum amyloid A
Muscle Specific Signature		PC 1	
X00371	545	0.3348	myoglobin
M33772	1527.7	0.3348	fast skeletal muscle troponin C
Z20656	2992.5	0.3083	cardiac alpha-myosin heavy chain
M21494	410.4	0.287	muscle creatine kinase (CKMM)
U96094	363.6	0.2803	sarcolipin (SLN)
J04760	701.8	0.279	slow-twitch skeletal troponin I (TNN1)
M83308	5723.7	0.2651	mitochondrial cytochrome-c oxidase subunit VIa
10103300	3/23./	0.2031	(COX6A)
X06825	452.3	0.2444	skeletal beta-tropomyosin
L21715	851.7	0.2257	troponin I fast-twitch isom
M21665	488.5	0.2184	beta-myosin heavy chain
M19309	1149.9	0.2099	slow skeletal muscle troponin T, clone H22h
X90568	3169.9	0.2077	titin prot (clone hh1-hh54)
S73840	350.5	0.2022	type IIx myosin heavy chain
M20543	993.2	0.1917	skeletal alpha-actin
X16504	1016.3	0.168	X51957 and others
M20642	747.2	0.15	alkali myosin light chain 1
U35637	386.9	0.1345	nebulin/U35637
M29458	564.4	0.1056	carbonic anhydrase III
M86407	759.1	0.0813	alpha actinin 3 (ACTN3)
Brain Specific Signature		PC 2	
S72043	90.4306	0.4026	GIF=growth inhibitory factor
M13577	686.2963	0.4020	myelin basic prot (MBP)
S40719	20.5566	0.3366	glial fibrillary acidic prot
HG1877-HT1917	82.2133	0.2733	Myelin Basic prot
X99076	49.5985	0.1778	NRGN
U48437	23.3006	0.1033	amyloid precursor-like prot 1
J04615		0.1404	amyloid precursor-like prot i
J04013	5 0026	0.1202	lunus autoantigan (small nuclear ribanuclenanrat
	5.9926	0.1292	lupus autoantigen (small nuclear ribonuclepoprot snRNP SM-D)
D21267	5.9926 184.849	0.1252	snRNP SM-D) highly expressed prot
D21267 L07807	184.849 30.2311	0.1252 0.1162	snRNP SM-D) highly expressed prot dynamin
	184.849 30.2311 27.4526	0.1252 0.1162 0.1159	snRNP SM-D) highly expressed prot
L07807 HG3437-HT3628 L10373	184.849 30.2311 27.4526 18.2544	0.1252 0.1162 0.1159 0.1123	snRNP SM-D) highly expressed prot dynamin Myelin Proteolipid prot (clone CCG-B7) sequence
L07807 HG3437-HT3628 L10373 M16364	184.849 30.2311 27.4526 18.2544 9.3301	0.1252 0.1162 0.1159 0.1123 0.1071	snRNP SM-D) highly expressed prot dynamin Myelin Proteolipid prot (clone CCG-B7) sequence creatine kinase-B
L07807 HG3437-HT3628 L10373	184.849 30.2311 27.4526 18.2544 9.3301 3.7109	0.1252 0.1162 0.1159 0.1123 0.1071 0.0912	snRNP SM-D) highly expressed prot dynamin Myelin Proteolipid prot (clone CCG-B7) sequence creatine kinase-B prostaglandin D2 synthase
L07807 HG3437-HT3628 L10373 M16364 M98539 U44839	184.849 30.2311 27.4526 18.2544 9.3301	0.1252 0.1162 0.1159 0.1123 0.1071	snRNP SM-D) highly expressed prot dynamin Myelin Proteolipid prot (clone CCG-B7) sequence creatine kinase-B
L07807 HG3437-HT3628 L10373 M16364 M98539	184.849 30.2311 27.4526 18.2544 9.3301 3.7109	0.1252 0.1162 0.1159 0.1123 0.1071 0.0912	snRNP SM-D) highly expressed prot dynamin Myelin Proteolipid prot (clone CCG-B7) sequence creatine kinase-B prostaglandin D2 synthase
L07807 HG3437-HT3628 L10373 M16364 M98539 U44839	184.849 30.2311 27.4526 18.2544 9.3301 3.7109 3.1469	0.1252 0.1162 0.1159 0.1123 0.1071 0.0912 0.089	snRNP SM-D) highly expressed prot dynamin Myelin Proteolipid prot (clone CCG-B7) sequence creatine kinase-B prostaglandin D2 synthase putative ubiquitin C-terminal hydrolase (UHX1) unc-18 homolog unknown prot
L07807 HG3437-HT3628 L10373 M16364 M98539 U44839 D63851	184.849 30.2311 27.4526 18.2544 9.3301 3.7109 3.1469 10.9002	0.1252 0.1162 0.1159 0.1123 0.1071 0.0912 0.089 0.0863	snRNP SM-D) highly expressed prot dynamin Myelin Proteolipid prot (clone CCG-B7) sequence creatine kinase-B prostaglandin D2 synthase putative ubiquitin C-terminal hydrolase (UHX1) unc-18 homolog
L07807 HG3437-HT3628 L10373 M16364 M98539 U44839 D63851 Y09836	184.849 30.2311 27.4526 18.2544 9.3301 3.7109 3.1469 10.9002 16.17	0.1252 0.1162 0.1159 0.1123 0.1071 0.0912 0.089 0.0863 0.0838	snRNP SM-D) highly expressed prot dynamin Myelin Proteolipid prot (clone CCG-B7) sequence creatine kinase-B prostaglandin D2 synthase putative ubiquitin C-terminal hydrolase (UHX1) unc-18 homolog unknown prot Na+,K+-ATPase catalytic subunit alpha-III isoform neuronal growth prot 43 (GAP-43)
L07807 HG3437-HT3628 L10373 M16364 M98539 U44839 D63851 Y09836 M37457	184.849 30.2311 27.4526 18.2544 9.3301 3.7109 3.1469 10.9002 16.17 9.0757	0.1252 0.1162 0.1159 0.1123 0.1071 0.0912 0.089 0.0863 0.0838 0.0805	snRNP SM-D) highly expressed prot dynamin Myelin Proteolipid prot (clone CCG-B7) sequence creatine kinase-B prostaglandin D2 synthase putative ubiquitin C-terminal hydrolase (UHX1) unc-18 homolog unknown prot Na+,K+-ATPase catalytic subunit alpha-III isoform
L07807 HG3437-HT3628 L10373 M16364 M98539 U44839 D63851 Y09836 M37457 M25667	184.849 30.2311 27.4526 18.2544 9.3301 3.7109 3.1469 10.9002 16.17 9.0757 27.35	0.1252 0.1162 0.1159 0.1123 0.1071 0.0912 0.089 0.0863 0.0838 0.0805 0.0779	snRNP SM-D) highly expressed prot dynamin Myelin Proteolipid prot (clone CCG-B7) sequence creatine kinase-B prostaglandin D2 synthase putative ubiquitin C-terminal hydrolase (UHX1) unc-18 homolog unknown prot Na+,K+-ATPase catalytic subunit alpha-III isoform neuronal growth prot 43 (GAP-43)
L07807 HG3437-HT3628 L10373 M16364 M98539 U44839 D63851 Y09836 M37457 M25667 D78577	184.849 30.2311 27.4526 18.2544 9.3301 3.7109 3.1469 10.9002 16.17 9.0757 27.35 6.3676	0.1252 0.1162 0.1159 0.1123 0.1071 0.0912 0.089 0.0863 0.0838 0.0805 0.0779 0.0779	snRNP SM-D) highly expressed prot dynamin Myelin Proteolipid prot (clone CCG-B7) sequence creatine kinase-B prostaglandin D2 synthase putative ubiquitin C-terminal hydrolase (UHX1) unc-18 homolog unknown prot Na+,K+-ATPase catalytic subunit alpha-III isoform neuronal growth prot 43 (GAP-43) DNA for 14-3-3 prot eta chain
L07807 HG3437-HT3628 L10373 M16364 M98539 U44839 D63851 Y09836 M37457 M25667 D78577 L20814	184.849 30.2311 27.4526 18.2544 9.3301 3.7109 3.1469 10.9002 16.17 9.0757 27.35 6.3676 68.1413	0.1252 0.1162 0.1159 0.1123 0.1071 0.0912 0.089 0.0863 0.0838 0.0805 0.0779 0.0779	snRNP SM-D) highly expressed prot dynamin Myelin Proteolipid prot (clone CCG-B7) sequence creatine kinase-B prostaglandin D2 synthase putative ubiquitin C-terminal hydrolase (UHX1) unc-18 homolog unknown prot Na+,K+-ATPase catalytic subunit alpha-III isoform neuronal growth prot 43 (GAP-43) DNA for 14-3-3 prot eta chain glutamate receptor 2 (HBGR2)
L07807 HG3437-HT3628 L10373 M16364 M98539 U44839 D63851 Y09836 M37457 M25667 D78577 L20814 J04046	184.849 30.2311 27.4526 18.2544 9.3301 3.7109 3.1469 10.9002 16.17 9.0757 27.35 6.3676 68.1413 6.4909	0.1252 0.1162 0.1159 0.1123 0.1071 0.0912 0.089 0.0863 0.0838 0.0805 0.0779 0.0779 0.0735 0.0729	snRNP SM-D) highly expressed prot dynamin Myelin Proteolipid prot (clone CCG-B7) sequence creatine kinase-B prostaglandin D2 synthase putative ubiquitin C-terminal hydrolase (UHX1) unc-18 homolog unknown prot Na+,K+-ATPase catalytic subunit alpha-III isoform neuronal growth prot 43 (GAP-43) DNA for 14-3-3 prot eta chain glutamate receptor 2 (HBGR2) calmodulin

D82343	140.3644	0.0649	AMY
S82024	47.6237	0.06	SCG10=neuron-specific growth-associated
			prot/stathmin homolog
D49958	29.5755	0.0571	membrane glycoprot M6
M65066	15.0292	0.0541	cAMP-dependent prot kinase regulatory subunit RI-
			beta
D87465	9.7149	0.0532	KIAA0275
X86809	4.3215	0.0524	major astrocytic phosphoprot PEA-15

#### Validation of gene expression patterns using new samples

Additional samples (three each) from liver, muscle, and brain were collected in a subsequent experiment, profiled transcriptionally and analyzed by applying the above projection methods. Fig. 2b shows the projections of the gene expression data of the new liver samples using the loadings obtained from the projection of the genes in structure A (that discriminated the two liver samples from the remaining tissues in the initial data set). All three liver samples are clearly separated along the first PC from the non-liver tissues in the initial data set, underscoring the tissue-specific nature of these genes, and hinting at the construction of a "liver axis" along the first PC. The genes distinguishing liver from non-liver tissues include albumin and those associated with the coagulation pathway (e.g. factor IX, antithrombin III and heparin cofactor), complement pathway (e.g. C8), lipid process (e.g. apolipoproteins), bile metabolism (e.g. fatty acid binding protein 1), xenobiotic metabolism (e.g. cytochrome P450), and iron homeostasis (e.g. hemopexin), a result which is to be expected based on the known biology of the liver. An examination of the 24 genes in this structure revealed that 33% of all gene pairs had correlation coefficients greater than 0.88 for these five liver samples. This value of the coefficient is significant at the 95% statistical level. Thus, a subset of these genes are expressed proportionately to each other in the liver tissue. For instance, it is known that apolipoprot H binds to negatively charged heparin, and the heparin cofactor and antithrombin III are serine proteases that inhibit the coagulation pathway (McNally et al., 1994, Vander et al., 1994).

The loadings of the 19 genes in structure B were similarly used to project the three new skeletal muscle samples, and the results are shown in Fig. 2c. Similar to the liver samples, the first PC clearly separates the new skeletal muscle samples, and acts like a "muscle axis". The genes include those associated with the cytoskeleton (e.g. actin,  $\alpha$ 1, actinin  $\alpha$ 3, nebulin), contraction (e.g. tropomyosin, troponin, myosin), glucose metabolism (e.g. enolase 3 $\beta$ ), CO2 metabolism (e.g. carbonic anhydrase III), and energy transduction (e.g. creatine kinase). Particularly, actinin  $\alpha$ 3 is known to have expression limited to skeletal muscle (North *et al.*, 1999), and carbonic anhydrase III is strictly present at high levels in skeletal muscle and much lower levels in cardiac and smooth muscle (Lloyd *et al.*, 1986). About 74 % of all gene pairs, after discounting ones with the same genes, had a correlation coefficient greater than 0.811, the 95% confidence level with the given number of samples. This rather striking degree of linear correlation implies that these genes are expressed proportionately in skeletal muscle samples, and may be coordinately regulated. For example, while both actin and myosin provide force

for muscle contraction, troponin, a regulatory protein, prevents actin and myosin interaction in resting muscle tissue. And, tropomyosin, an actin filament binding protein is required for the interaction of actin and troponin. It is also known that titin maintains resting tension in skeletal muscle (Vander *et al.*, 1994).

Finally, the 86 genes in structure C were used to project the new brain samples, and as Figure 2d shows, the new brain samples are clearly separated from the other non-brain samples, and fall in the same region as the brain samples of the initial set. The genes include those associated with myelin structure (e.g. myelin basic protein), astrocytic differentiation (e.g. glial fibrillary acidic protein), synaptic reorganization (e.g. calmodulin, neurogranin, and GAP-43) and neurotransmission (e.g. glutamate receptor). Of note, many genes with no known functions are also reported here to be specific for the brain samples.

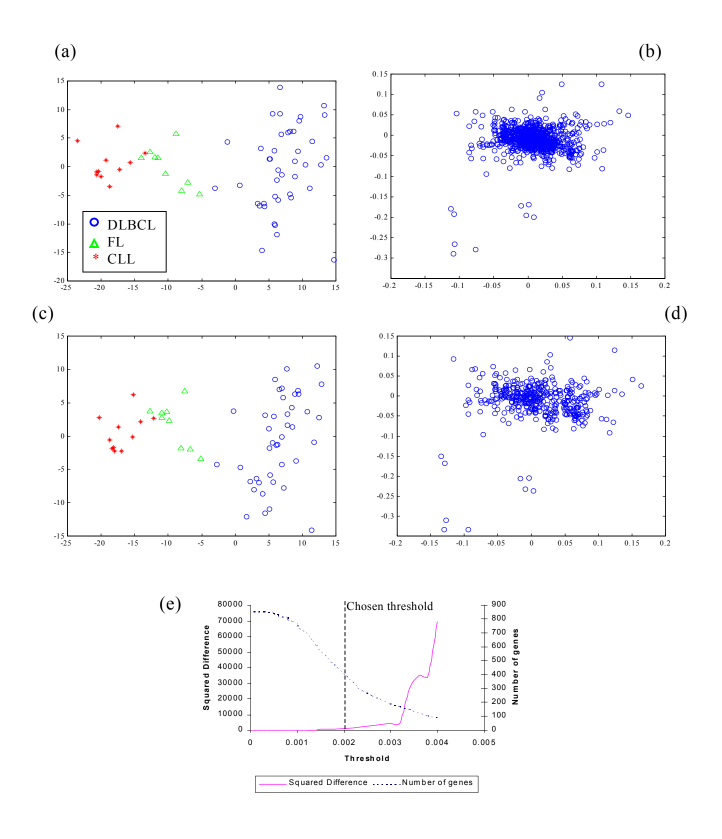
The use of projection methods to analyze the effect of these genes on the samples also led to the automatic construction of a reduced dimension classifier space for the liver, muscle, and brain tissues. As demonstrated here, new samples may be projected onto this space, and the score value used to classify the tissue sample.

#### **Application to other data sets**

Figure 3 shows the result of the application of the current methodology to the gene expression data on lymphoid malignancies (Alizadeh *et al.*, 2000). Expression phenotype of 62 samples of diffuse large B-cell lymphoma (DLBCL), follicular lymphoma (FL), and chronic lymphotic leukemia (CLL) were measured on 17,856 cDNA clones. A simple projection reveals the presence of two clusters, and one intervening group of samples. Querying the nature of these samples reveals an almost perfect segmentation of the samples, in a PC space that comprises a mere 35% of the information in the data. Implementing the thresholding procedure allows for the identification of 401 consequential genes, which maintain the patterns in the data, with minimal distortion. No outstanding structures suggest themselves in the loading plot. The observation of linear structures is a unique characteristic of each data set, and will not necessarily occur in all cases. In this particular case, just the thresholding procedure is sufficient to allow for segmentation of the samples and identification of consequential genes.

#### DISCUSSION

We have demonstrated the utility of PCA as an initial step in the analysis of microarray data to extract and examine gene expression patterns. Previous work has applied a similar approach (singular value decomposition) to construct linear combinations of gene expressions (called "characteristic modes" or "eigengenes") from microarray measurements of time-series samples (Alter *et al.*, 2000, Holter *et al.*, 2000). Here, we extend the application of PCA to the analysis of non-time series data, and the data-driven



<u>Figure 3:</u> Projection of the Lymphoma samples using the PCA. The projection already reveals a fairly clear separation of the three classes in the data. The thresholding procedure allows for the identification of 401 genes from about 850 cDNA clones which are sufficient to describe the patterns observed. (a) and (b) are the score and the loading plot prior to thresholding and (c) and (d) are the score and loading plot post thresholding. (e) reveals the effect of thresholding on the number of genes retained and the squared difference. The chosen threshold, 0.002 is the point beyond which the squared difference explodes.

learning and sample classification problem. The reason for the broad applicability of the PCA lies in its strong, yet flexible, mathematical structure, and the correspondence between the score plot and the loading plot. This latter feature is exploited in the interactive methodology presented for the elimination of redundant variables or genes. This method is general, and may be applied to any data set.

Our methodology facilitated the identification of strong underlying structures in the data. The identification of such structures is uniquely dependent on the data, and is not generally guaranteed. For example, the expression data on leukemia samples (Golub *et al.*, 1999) was similarly analyzed, however, no evident patterns presented themselves, though diffuse structures containing some discriminatory information could be observed at higher, less informative PCs (data not shown). This may be due to the fact that the PCA attempts to maximize the variation that it captures in the data. In cases where the discriminatory information is not the most important type of variation (perhaps due to the presence of a large number of non-discriminatory genes), the above analysis will not yield discriminatory patterns between two classes of tissues/sample. When discriminatory genes are preselected by applying a t-test on pre-classified samples, and used for projection, clear separations are obtained between AML and ALL classes.

Several genes in the tissue-specific signatures identified here are justifiable with respect to known biology regarding the particular tissue. In the case of the liver and muscle samples, co-ordinate expression of some of these genes may also be biologically explained. Elucidation of the function and role of the other genes observed in these tissue-specific signatures must await further experiments.

In the current study, the data was not mean-centered. Mean-centering is geometrically equivalent to shifting the origin of the PCA coordinate system to the centroid of the data, a procedure which may or may not yield different results. For the purposes of comparison, the data was mean-centered, and then analyzed as described above. The structures for the liver and muscle samples were identified in the first and second PC, while the identification of the brain structure required the inclusion of the third PC. The list of genes identified overlapped strongly with the one presented here. This raises our confidence in the significance of the genes identified, but also underscores the fact that different processing methods will give rise to a slightly different list of genes, and it may be best to adopt several processing methods, and choose a common subset of genes.

Projection methods shift the focus of analysis from individual genes to the combined quantitative effect of several consequential genes. Here, due to the strong structures observed in the data, such a combination led to the construction of reduced dimension classifiers for the liver, muscle and brain tissues. If the sole objective of the analysis is to yield a classifier, then other projection methods, such as Canonical Discriminant Analysis (Hwang *et al.*, in preparation) are more appropriate, and rigorous. If the objective is data exploration, the PCA is better applied, since few *a priori* assumptions, such as sample class type, are made. Overall, due to their data reduction properties and their flexibility in dealing with large data sets, projection methods are an important class of tools for the analysis of microarray data.

#### **METHODS**

#### Data treatment

Each array from the BWH data was scaled to a target intensity of 100. All negative expression values were reduced to zero for the purposes of analysis. For treatment of the lymphoma data, refer Alizadeh *et al.* (2000). In the Lymphoma data set, genes which have missing values for the 62 experiments are removed from the analysis. This leaves us with an initial starting number of 854 cDNA clones.

#### Principal Components Analysis

Singular value decomposition is used to calculate the principal components of a data matrix (Dillon and Goldstein, 1984). Any data matrix X with S samples (tissues) on the rows and V variables (genes) on the columns may be decomposed as follows:

$$X = U T L'$$

$$(sxv) = (sxR)(RxR)(Rxv)$$
(1)

where T is a diagonal matrix whose values are the singular values of the matrix X. The singular values of X are the square roots of the nonzero eigenvalues of the square matrix X'X as well as XX' (X' being the transpose of X). The columns of U and L contain the eigenvectors of XX' and X'X, respectively. The maximum number of independent dimensions, R is determined by the rank of the matrix X.

The *loadings* of the genes, or their coefficients in the linear combination which forms the principal component, is given by the column vectors of the matrix L. The magnitude of a gene loading is a measure of its importance in defining the principal component. The *scores* of the samples, or the *projections* of the samples on the principal components is given by:

$$Sc = X L$$
 (2)

The amount of information in the data that the first r principal components capture may be quantified as:

% information captured by the first 
$$r$$
 components (out of R total) = 
$$\frac{\sum_{i=1}^{r} SV_i^2}{\sum_{i=1}^{R} SV_i^2}$$
 (3)

where SV<sub>i</sub> is the ith singular value.

The filter on the loadings was implemented by dividing each loading by the sum of the magnitudes of all the other loadings for that PC, and then rejecting all genes with a loading less than the threshold value. The distortion of patterns in the score plot due to the removal of genes in this thresholding procedure was measured by the sum of the squares of the difference between the 40 original score values and the 40 score values produced with the filtered gene set. Mathematically,

$$SD = \sum_{s=1}^{40} \sum_{i=1}^{5} (y_{s,i,f} - y_{s,i,o})^2$$
 (4)

where SD is the squared difference,  $y_{s,i,o}$  is the score value of the  $s^{th}$  sample on the  $i^{th}$  PC in the projection using all the 7070 genes, while  $y_{s,i,f}$  is the score value of the  $s^{th}$  sample on the  $i^{th}$  PC obtained when a filtered gene set is used.

Gene lists and supplementary materials

All relevant gene sets and supplementary figures will be posted as web-site supplements at the Genome Research site.

#### **ACKNOWLEDGEMENTS**

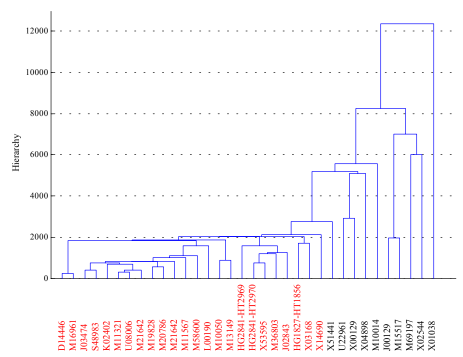
We would like to acknowledge the anonymous reviewers for their constructive suggestions for the paper. This work was supported by a grant from the Engineering Research Program of the Office of Basic Energy Science at the Dept. of Energy, Grant No. DE-FG02-94ER-14487 and DE-FG02-99ER-15015, and NIH Grant No. 1-RO1-DK58533-01.

#### REFERENCES

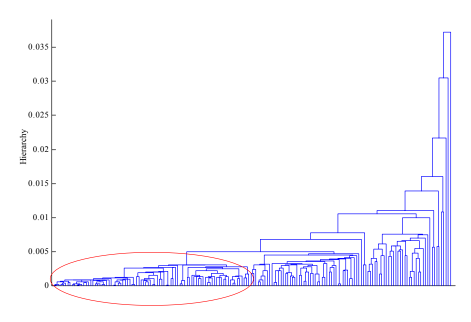
- 1. Alizadeh, A. A., Eisen, M. B., Davis, R. E., Ma, C., Lossos, I. S., Rosenwald, A., Boldrick, J. G., Sabet, H., Tran, T., Yu, X., *et al.* 2000. Distinct types of diffuse large B-cell lymphoma identified by gene expression profiling. *Nature*, **403**, 503-511.
- 2. Alter, O., Brown, P. O., and Botstein, D. 2000. Singular value decomposition for genome-wide expression data processing and modeling. *Proceedings Of the National Academy Of Sciences Of the United States Of America*, **97**, 10101-10106.
- 3. Brown, M. P. S., Grundy, W. N., Lin, D., Cristianini, N., Sugnet, C. W., Furey, T. S., Ares, M., and Haussler, D. 2000. Knowledge-based analysis of microarray gene expression data by using support vector machines. *Proceedings Of the National Academy Of Sciences Of the United States Of America*, 97, 262-267.
- 4. Dillon, W. R. and Goldstein, M. 1984. *Multivariate Analysis*. John Wiley & Sons, USA.
- 5. Eisen, M. B., Spellman, P. T., Brown, P. O., and Botstein, D. 1998. Cluster analysis and display of genome-wide expression patterns. *Proceedings Of the National Academy Of Sciences Of the United States Of America*, **95**, 14863-14868.
- 6. Golub, T. R., Slonim, D. K., Tamayo, P., Huard, C., Gaasenbeek, M., Mesirov, J. P., Coller, H., Loh, M. L., Downing, J. R., Caligiuri, M. A., Bloomfield, C. D., and Lander, E. S. 1999. Molecular classification of cancer: Class discovery and class prediction by gene expression monitoring. *Science*, **286**, 531-537.
- 7. Holter, N. S., Mitra, M., Maritan, A., Cieplak, M., Banavar, J. R., and Fedoroff, N. V. 2000. Fundamental patterns underlying gene expression profiles: Simplicity from complexity. *Proceedings Of the National Academy Of Sciences Of the United States Of America*, **97**, 8409-8414.

- 8. Hsiao, L., Dangond, F., Yoshida, T., Hong, R., Jensen, R.V., Misra, J., Dillon, W., Lee, K., Clark, K., Haverty, P., *et al.* 2001. A compendium of gene expression in normal human tissues reveals tissue-selective genes and distinct expression patterns of housekeeping genes. *Physiological Genomics* (in press).
- 9. Hughes, T. R., Marton, M. J., Jones, A. R., Roberts, C. J., Stoughton, R., Armour, C. D., Bennett, H. A., Coffey, E., Dai, H. Y., He, Y. D. D., *et al.* 2000. Functional discovery via a compendium of expression profiles. *Cell*, **102**, 109-126.
- 10. Kamimura, R. T. 1997. Application of Multivariate Statistics to Fermentation Database Mining. Ph.D. Thesis, Department of Chemical Engineering, Massachusetts Institute of Technology.
- 11. Lloyd J., McMillan, S., Hopkinson, D., and Edwards, Y.H. 1986. Nucleotide sequence and derived amino acid sequence of a cDNA encoding human muscle carbonic anhydrase. *Gene*, **41**, 233-239.
- 12. McNally, T., Cotterell, S. E., Mackie, I. J., Isenberg, D. A., and Machin, S. J. 1994. The Interaction of Beta(2) Glycoprotein-I and Heparin and Its Effect on Beta(2) Glycoprotein-I Antiphospholipid Antibody Cofactor Function in Plasma. *Thrombosis and Haemostasis*, 72, 578-581.
- 13. North, K. N., Yang, N., Wattanasirichaigoon, D., Mills, M., Easteal, S., Beggs, A. H. 1999. A common nonsense mutation results in alpha-actinin-3 deficiency in the general population. *Nature Genetics*, **21**, 353-354.
- 14. Perou, C. M., Jeffrey S. S., Van de Rijn, M., Rees, C. A., Eisen, M. B., Ross, D. T., Pergamenschikov, A., Williams, C. F., Zhu, S. X., Lee, J. C. F., Lashkari, D., Shalon, D., Brown, P. O., and Botstein, D. 1999. Distinctive gene expression patterns in human mammary epithelial cells and breast cancers. *Proceedings Of the National Academy Of Sciences Of the United States Of America*, **96**, 9212-9217.
- 15. Rannar S., MacGregor, J.F, Wold, S. 1998. Adaptive Batch Monitoring using Hierarchical PCA. *Chemometrics & Intelligent Laboratory Systems*, **41**, 73-81.
- 16. Spellman, P. T., Sherlock, G., Zhang, M. Q., Iyer, V. R., Anders, K., Eisen, M. B., Brown, P. O., Botstein, D., and Futcher, B. 1998. Comprehensive identification of cell cycle-regulated genes of the yeast Saccharomyces cerevisiae by microarray hybridization. *Molecular Biology Of the Cell*, **9**, 3273-3297.
- 17. Tamayo, P., Slonim, D., Mesirov, J., Zhu, Q., Kitareewan, S., Dmitrovsky, E., Lander, E. S., and. Golub, T. R. 1999. Interpreting patterns of gene expression with self-organizing maps: Methods and application to hematopoietic differentiation. *Proceedings Of the National Academy Of Sciences Of the United States Of America*, **96**, 2907-2912.
- 18. Vander, A. J., Sherman, J. H., and Luciano, D. H. 1994. *Human Physiology*. McGraw-Hill, NY.

#### SUPPLEMENTARY FIGURES



**Supplementary Figure 1**: Clustering of the genes identified in structure A to identify genes with the same angle, but far removed from the actual structure. Single linkage clustering with a euclidean distance metric was used for clustering the samples. Of the 34 genes in the structure, the 24 genes in red were the ones chosen for subsequent analysis and projections, since they cluster tightly together, while the others are further away.



**Supplementary Figure 2**: Clustering of the genes identified in structure C to identify genes with the same angle, but far removed from the actual structure. Single linkage clustering with a euclidean distance metric was used for clustering the samples. Of the 174 genes in the structure, the 86 genes enclosed by the red circle were the ones chosen for subsequent analysis and projections, since they cluster tightly together, while the others are further away.