## PCA and plotting:

Scree plot: eigenvalues in non-increasing order



2D plot of the data cloud projected on the plane spanned by the first two principal components; this captures more variability than any other 2D projection of the cloud



3D plot of the data cloud projected on the space spanned by the first three principal components; this captures more variability than any other 3D projection of the cloud

Cannot go further up and maintaining a complete view...

But we can use a scatter plot matrix; 2D side views of the T-dimensional cloud. This can be done with the original coordinates as well. When done with the principal components, the choice of side views is guided by variability explanation



Each plot in the matrix is usually rescaled to optimal "filling"

*S* is very sensitive to anomalous and/or "out of the bulk points.

In the 2D plot of the first two principal components, we might detect points that inflate variances and/or distort covariances (influential on the main variability directions).

In the 2D plot of the last two principal components, we might detect points that induce directions that are not really there; mask singularity (influential on the small variability directions, and how many directions one chooses to retain).

Main <u>variability axes may be different</u>, and <u>smaller variability axes may not exist</u>, without these points. Need to identify and investigate them and attempt to understand if their anomalous position is due to errors.

Try analysis with and without.

Use a "robust version" of S.



### Second phase of PCA: deciding how many directions to retain (K)

e.g. Is the first plane enough? We reason in terms of variability, and we'll decide based on it, analyzing the eigenvalues – whether it is enough with respect to other structural features is a different issue!

1. Consider the proportion of explained variability, and retain as many directions as needed to explain a selected proportion



3. Consider the average explained variability per component, and retain directions with an explanatory capability above average – on the scree plot:



4. Look for bends in the scree plot. If there is a clear bend, keep directions associated with eigenvalues before the bend – those afterwards have comparable, small(er) size (smaller the more they are)



5. "Testing" version of 3. *If the data is elliptical (Gaussian or about)*, we can perform a sequence of tests to assess how many tail eigenvalues are statistically equal to one another. This is based on a large N chi-square null distribution (not clear whether normality is a viable assumption)

$$H_{0}: \quad \lambda_{T-h+1} = \dots = \lambda_{T}$$

$$H_{1}: \quad not$$

$$\left(N - \frac{2T - 11}{6}\right) \left(h \ln \overline{\lambda} - \sum_{j=T-h+1}^{T} \ln \lambda_{j}\right) \quad \stackrel{approx}{\sim} \quad \chi_{\frac{1}{2}(h-1)(h+2)}^{2}$$

More sophisticated methods exist – fairly large literature.

## What are we missing?

With 1, we fix the percentage of variability we are missing

With 2,3 and 4 we don't. It might not be negligible

With 1 and 2, there might still be variability structure in the neglected directions

With 3 and 4, the variability is approximately the same in all neglected directions

If the percentage of variability we are missing is very small (e.g. 2%), we might argue that whatever structural feature might live in the neglected directions, it occurs on a scale so small that we do not care.

But if the percentage of variability we are missing is not very small (e.g. 20 or 30%), we ought to investigate what is going on there.



Of course if we are neglecting more than two or three directions a direct graphical investigation will not be possible.

# Think about a "scrambling scheme" to provide a chance background for the choice of K ("reference curves" on a scree plot).

#### **Interpreting principal components:**



Using principal components to rank (and select) genes:

$$X_i = \sum_{j=1}^T W_{i,j} V_j$$

How close is  $X_i$  to  $V_j$ ?

$$P_{V_{j}}X_{i} = (W_{i,j})V_{j}$$

$$\frac{\|P_{V_{j}}X_{i}\|^{2}}{\|X_{i}\|^{2}} = \frac{(W_{i,j})^{2}}{\sum_{j=1}^{T} (W_{i,j})^{2}} = corr^{2}(X_{i},V_{j}) = R_{X_{i}|V_{j}}^{2}$$

(recall one norm is =1)

How close is  $X_i$  to  $V_1 \dots V_K$  as a group, i.e. to  $Span(V_1 \dots V_K)$ ?

$$P_{Span}X_{i} = \sum_{j=1}^{K} P_{V_{j}}X_{i} = \sum_{j=1}^{K} (W_{i,j})V_{j}$$

$$\frac{\|P_{Span}X_{i}\|^{2}}{\|X_{i}\|^{2}} = \frac{\sum_{j=1}^{K} \|P_{V_{j}}X_{i}\|^{2}}{\|X_{i}\|^{2}} = \frac{\sum_{j=1}^{K} (W_{i,j})^{2}}{\sum_{j=1}^{T} (W_{i,j})^{2}} = R_{X_{i}|V_{j}...V_{K}}^{2}$$

Determination coefficient from a linear ols fit (ith gene on K pca components, using the T conditions... rows)

Genes closest to the natural direction(s) of highest variability in expression Gene profiles better reconstructed in terms of main expression patterns:



<u>Think about a "scrambling scheme" to provide a chance background for the</u> ranking ("reference curves" on the ranking plot).

### Using the correlation matrix instead of the covariance matrix:

Essential if the original coordinates represent quantities with different units of measure (not the case for us).

Also important if the original coordinates have very different variability sizes, and we do not want to take these differences under consideration... we are after "standardized variability structure"; that is, <u>correlation structure</u> (for us, think of a situation in which expression variability is much higher in some conditions than in others, and we do not consider these differences as relevant for our purposes).

$$S = \frac{1}{N} \sum_{i=1}^{N} (X_i - \overline{X}) (X_i - \overline{X})' = \begin{pmatrix} s_1^2 & s_{1,T} \\ & \ddots & \\ s_{T,1} & s_T^2 \end{pmatrix}$$

$$R = D(1/s_j)SD(1/s_j) = \frac{1}{N} \sum_{i=1}^{N} D(1/s_j)(X_i - \overline{X})(X_i - \overline{X})'D(1/s_j) = \begin{pmatrix} 1 & r_{1,T} \\ & \ddots & \\ r_{T,1} & 1 \end{pmatrix}$$

Correlation matrix, i.e. var/cov matrix of the standardized cloud  $D(1/s_j)(X_i - \overline{X})$ 

We are standardizing the data matrix by column.

### The eigenstructures of S and R are different in both eigenvectors and eigenvalues.

References:

Gnanadesikan: Methods for Statistical Data Analysis of Multivariate Observations. Wiley.

Neal S. Holter, Madhusmita Mitra, Amos Maritan, Marek Cieplak, Jayanth R. Banavar, and Nina V. Fedoroff: Fundamental patterns underlying gene expression profiles: Simplicity from complexity PNAS 97: 8409-8414;

Orly Alter, Patrick O. Brown, and David Botstein: Singular value decomposition for genome-wide expression data processing and modeling PNAS 97: 10101-10106.

(Important: Singular Value Decomposition and PCA are equivalent analyses).

Read these paper and match their logic in, and use of, dimension reduction with the considerations we have made.

- How does "noise" look like?
- How do we construct a reference chance scenario?
- Under what circumstances can we claim that something "structured" we are throwing away is "non-experimental"? ("experimental artifacts")

Also, start thinking of the use of dimension reduction in connection with clustering techniques (PCA is <u>NOT</u> a clustering technique).