# Module 2: Introduction to Statistics 

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## Topic

- Multiple Testing
- Family-Wise Error Rate
- Bonferroni adjustment
- False Discovery Rate
- Multivariate Data Analysis
- Principle Component
- Screen plot, Bipolar plot
- Cluster Analysis: K-means, Hierarchical Clustering
- Dendrograms
- Heatmaps


## Multiple Testing

- Multiple testing is frequently used in large data sets, particularly in discovery science
- E.g. Genome-wide association studies (GWAS) - test up to several million genetic variants for association with a trait
- When many tests are performed the following questions are relevant:
- Are there any true positive results?
- How many are false positive?
- Which are the true positives?


## Type I Error: False Positive (FP) and True Positive

 (TP) Rate- Type I error or False Positive Rate for testing a single hypothesis usually is set at $\alpha=0.05$

$$
F P=\operatorname{Pr}\left(\text { Reject } H_{0} \mid H_{0} \text { is true }\right)=.05
$$

$$
T N=\operatorname{Pr}\left(\text { Do not Reject } H_{0} \mid H_{0} \text { is true }\right)=.95(=1-\mathrm{FP})
$$

- Suppose we are testing two independent null hypotheses $H_{01}$ and $H_{02}$. In such case the Type I Error is compounded

$$
\operatorname{Pr}\left(\text { Reject } H_{01} \text { or } H_{02} \mid H_{01} \text { and } H_{02} \text { are true }\right)=1-(.95)^{2}=.0975
$$

- In general, the more hypotheses you test, the more likely it is to see by chance a difference that is not there


## Family-Wise Error Rate (FWER)

- Suppose we test a family of hypotheses, e.g. we collected 10 possible biomarkers for high BP, and test whether:
- Marker 1 does (or does not $H_{01}$ ) relate with high BP
- Marker 2 does (or does not $H_{02}$ ) relate with high BP
- Marker m.......
- If we perform all the tests, and they are independent, the probability that we make at least one false positive (or "false discovery" $)$ is around $\underline{0.4}\left(=1-(.95)^{10}\right)$


## Family-Wise Error Rate (FWER)

- Similarly, if we test if a treatment effects 10 outcomes (BP, diabetes,...,lung cancer), the probability of making at least one false positive is still around $\underline{0.4}$
- This is called the "family-wise error rate" (FWER):
- FWER $=\operatorname{Pr}\left(\right.$ Reject at least one of $H_{0 k} \mid$ All $H_{0 k}$ are true)
- FWER is always greater than $\alpha=0.05$ and could be quite large if number of tests, $m$, is large


## Controlling Family-Wise Error Rate (FWER): Bonferroni Adjustment

- Let $H_{0 k}$ be a family of $k=1,2, \ldots, m$ hypotheses. If we reject $H_{0 k}$ when $p_{k}<\alpha$, then the following is true:

$$
\begin{equation*}
\text { FWER }=\operatorname{Pr}\left(\text { Reject at least one of } H_{0 k} \mid A l l H_{0 k} \text { are true }\right) \leq m \alpha \tag{1}
\end{equation*}
$$

- From Eq. (1), if we carry out the significance of each test at $p_{k}<\alpha^{*}$ where $\alpha^{*}=\frac{\alpha}{m}$, then the FWER is at most:

$$
\mathrm{FWER} \leq m \alpha^{*}=m \frac{\alpha}{m}=\alpha
$$

## Controlling Family-Wise Error Rate (FWER): Bonferroni Adjustment

- Bonferroni adjustment: If $\alpha=0.05$ and there are $\mathrm{m}=10$ tests, then use $\frac{\alpha}{10}=.005$ as a criteria to reject a null hypothesis, i.e. $\mathrm{p}<.005$
- Bonferroni adjustment works OK for classical multiple testing (when $m$ ~ 3-5). But in general it is too conservative. It overprotects against FWER and, as a result, the Power is reduced.
- For a large number of multiple testing, the False Discovery Rate (FDR) method is a better alternative


## False Discovery Rate (FDR)

- FDR is the expected rate of false discoveries among all discoveries (rejected null hypotheses)

$$
F D R=\frac{\# \text { False Discoveries }}{\# \text { All Discoveries }}
$$

- E.g. If there were $m=1000$ discoveries ( 1000 null hypotheses were rejected) and a FDR level (q-value) for these tests was 0.05 , then 50 among 1000 discoveries were expected to be false discovery
- How to adjust for multiple testing so that FDR $\leq .05$ ?
- For each of the $m$ tests, get the p -value. Order them: $p_{1} \leq p_{2} \leq \ldots$ $\leq p_{m}$. Find the largest k , such that $p_{k} \leq \frac{k * .05}{m}$, then reject $H_{01}, \ldots, H_{0 k}$


# Illustration from the Example in Benjamini et. al. Article on FDR 

Benjamini, Yoav; Hochberg, Yosef (1995). Controlling the false discovery rate: a practical and powerful approach to multiple testing. Journal of the Royal Statistical Society, Series B (Methodological) 57 (1): 289-300.
http://www.math.tau.ac.il/~ybenja/MyPapers/benjamini hochberg1995.pdf

Example: 15 tests resulted in the following 15 p-values:
$0.0001,0.0004,0.0019,0.0095,0.0201,0.0278,0.0298,0.0344$, $0.0459,0.3240,0.4262,0.5719,0.6528,0.7590,1.000$.

## Example: Bonferroni and FDR Adjustment ( $\mathrm{m}=15$ tests)

$k \quad p_{k}$ (order) $\quad \alpha=.05$

| 1 | .0001 | $<.05$ |
| :--- | :---: | :---: |
| 2 | .0004 | $<.05$ |
| 3 | .0019 | $<.05$ |
| 4 | .0095 | $<.05$ |
| 5 | .0201 | $<.05$ |
| 6 | .0278 | $<.05$ |
| 7 | .0298 | $<.05$ |
| 8 | .0344 | $<.05$ |
| 9 | .0459 | $<.05$ |
| 10 | .3240 | $>.05$ |
|  | $\ldots$. | $\ldots$ |
| 15 | 1.000 | $>.05$ |

## Example: Bonferroni and FDR Adjustment

 ( $\mathrm{m}=15$ tests)$k \quad p_{k}$ (order) $\quad \alpha=.05$

| 1 | .0001 | $<.05$ |
| :--- | :--- | :--- |
| 2 | .0004 | $<.05$ |
| 3 | .0019 | $<.05$ |
| 4 | .0095 | $<.05$ |
| 5 | .0201 | $<.05$ |
| 6 | .0278 | $<.05$ |
| 7 | .0298 | $<.05$ |
| 8 | .0344 | $<.05$ |
| 9 | .0459 | $<.05$ |
| 10 | .3240 | $>.05$ |

15 $1.000>.05$

Bonferroni
$\alpha=\frac{.05}{m}=.0033$
<. 0033
<. 0033
<. 0033
$>.0033$
$>.0033$
...
...

$$
\begin{gathered}
\text { FDR( } q=.05) \\
\frac{k * .05}{m} \\
<.0033 \\
<.0066 \\
<.0099 \\
\leq .0132 \\
>.0165 \\
>.0198 \\
>.0231 \\
>.0264 \\
>.0297 \\
>.0330 \\
\ldots \\
>.05
\end{gathered}
$$

- FDR: $\underline{\mathrm{k}=4}$ is the largest $k$ for which $p_{k} \leq \frac{k * .05}{m}$. Thus, reject for $p_{1}, p_{2}, p_{3}, p_{4}$


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- False Discovery Rate
- Multivariate Data Analysis
- Principle Component
- Screen plot, Bipolar plot
- Cluster Analysis: K-means, Hierarchical Clustering
- Dendrograms
- Heatmaps


## Multivariate Analysis

- Multivariate analysis is different from the other modeling techniques (e.g. t-test, regressions) because there is no outcome or predictor
- In multivariate statistics we look for structure in the data
- Two common methods that look for structure are:
- Principal Component Analysis: Look for structure among variables
- Cluster Analysis: Look for structure among individuals


## Multivariate Analysis

- In multivariate data the number of variables of interest ( $X_{1}, X_{2}, \ldots, X_{p}$ ) may be large or too large (e.g. high dimensional data)
- This may cause problems with statistical modeling (i.e. regression) If $p>n$ then the degrees of freedom ( $d f=n-p-1$ ) for regression would be negative. In that case one can't run a multiple regression (Not enough data points (n) to estimate $p$ parameters)
- The interpretation of large data or results will be cumbersome
- There may be multiple testing issues (e.g. many false discoveries)
- Thus, data reduction when dealing with multivariate data is needed


## Principal Components Analysis (PCA)

- Principal component analysis (PCA) is a dimension-reduction method that generates a new set of decorrelated variables
- The new variables, called Principal Components (PC), are linear combinations of the original variables $\left(X_{1}, X_{2}, \ldots, X_{p}\right)$
- The idea of PCA is to find a small number of linear combinations of the variables (X's), which capture most of the variation of the original data


## Principal Components Analysis (PCA)

- Simple example: Suppose, that you had four measures (i.e. exam scores in math, biology, physics, chemistry). How would you summarize overall performance into a single score?
- A solution is to take the mean of the four variables

$$
S=\frac{x_{1}+x_{2}+x_{3}+x_{4}}{4}=\frac{1}{4} x_{1}+\frac{1}{4} x_{2}+\frac{1}{4} x_{3}+\frac{1}{4} x_{4}
$$

- S is a linear combination of $x_{1}, x_{2}, x_{3}, x_{4}$ with coefficient $/=(1 / 4,1 / 4,1 / 4,1 / 4)$
- PCA is statistical technique that finds few linear combinations (similar to S) that summarize the data


## Principal Component Analysis (PCA)

Original Data:

PC:

$-P C=l_{1} X_{1}+l_{2} X_{2}+\cdots+l_{p} X_{p}$
$-l_{1}, l_{2}, \ldots, l_{p}$ are called the loading factors for $P C$ (standardized so: $\sum l_{i}{ }^{2}=1$ ). They show how each X contributes to the $P C$

- PC's are uncorrelated: $\operatorname{Corr}\left(P C_{i}, P C_{j}\right)=0$ :
- $P C_{k}$ are ordered so the first one $\left(P C_{1}\right)$ explains most of the variance and so on


## Example: Principal Component Analysis (TROPHY Data)

- Part of the metabolic risk score can be measured using the following10 variables:
- Insulin
- Glucose
- Ins:Gluc ratio
- Triglycerides
- HDL
- LDL
- HDL:LDL ratio
- Total Cholesterol
- Systolic blood pressure
- Diastolic blood pressure
- Each measure represents a health risk (cardiovascular risk). For HDL and HDL:LDL low score is bad, for the rest high score is bad


## Example: Principal Component Analysis (TROPHY Data)

- So what happens when some scores are good and some are bad? We will use PCA to summarize the data in few meaningful PC's that still carry most of the information?
- Calculating principal components is easy (using R/SAS)
- Interpreting what the components mean in scientific terms is not always easy


## Example: Principal Component Analysis (TROPHY Data)

Output in R:

| Importance of components: | PC1 | PC2 | PC3 | PC4 | PC5 | PC6 | PC7 | PC8 |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Proportion of Variance | .29 | .22 | .13 | .12 | .09 | .08 | .05 | .01 |
| Cumulative Proportion | .29 | .51 | .65 | .77 | .86 | .94 | .99 | 1.0 |

- PC1 alone explains $29 \%$ of the variance from the original data
- PC2 alone explains $22 \%$ of the variance from the original data
- Cumulative: PC1 and PC2 jointly explain 51\% of the variance from the original data
- How to choose the number of PC?
- Use Eigenvalues, Screen Plot


## Screen Plot: Selecting the Number of PC's



- Eigenvalue criteria: Choose all PC's for which Eigenvalue > 1
- Visually: Look for the number of PC's where the curves start to flatten
- Explained Variance: Choose a small \# of meaningful PC's that explain a "sufficient" amount of variance (e.g. 50-60\%)


## Interpretation of PC's: TROPHY Data

- Importance of components: PC1 PC2 PC3 PC4 PC5 PC6 PC7 PC8

| Proportion of Variance | .29 | .22 | .13 | .12 | .09 | .08 | .05 | .01 |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Cumulative Proportion | .29 | .51 | .65 | .77 | .86 | .94 | .99 | 1.0 |

- What is the interpretation of PC1, PC2, PC3?


## Interpretation of PC's: TROPHY Data

 (Loading Factors)|  | PC1 | PC2 | PC3 |
| :--- | :---: | :---: | :---: |
| Insulin | 0.38 | 0.40 | -0.03 |
| Gluc | 0.25 | 0.13 | -0.04 |
| Ins_gluc | 0.38 | 0.39 | -0.05 |
| Trigl | 0.34 | -0.04 | -0.14 |
| HDL | -0.32 | -0.15 | 0.39 |
| LDL | 0.31 | -0.52 | 0.09 |
| hdl_Idl | -0.43 | 0.26 | 0.20 |
| Chol | 0.28 | -0.52 | 0.17 |
| SBP | 0.13 | 0.14 | 0.56 |
| DBP | 0.25 | 0.14 | 0.66 |

PC1=.38Ins+.25Gluc+.38Ins_gluc+.34Trig-.32HDL+.3LDL-.43HDL_LDL +.38Chol+.13SBP+.25DBP

## Biplot of PC1 vs. PC2 in SAS



## PCA Summary (Example: TROPHY Data)

- The metabolic risk profile based on the 10 measures in the TROPHY example can be summarized using 2-3 PC's: PC1, PC2, PC3
- PC1 is an overall weighted average score of the metabolic risk (high is bad)
- PC2 is a contrast score: Insulin - Lipids
- PC3 is an weighted average BP score (high is bad)
- PC1 and PC2 explain 51\% of the original variance
- PC1,PC2,PC3 explain 65\% of the original variance


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Multivariate Data Analysis

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- Cluster Analysis: K-means, Hierarchical Clustering
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- Heatmaps


## Cluster Analysis

- Cluster analysis is a set of techniques that look for groups (clusters) in the data such that:
- Individuals belonging to the same group resemble each other
- Individuals belonging to different groups are dissimilar
- There are two main approaches of carrying out such allocation
- Partitional: Partitioning into a number of clusters pre-specified by the user
- K-means Method
- Agglomerative: Starting with each individual as a separate cluster and aggregate similar individuals/clusters ending up with a single cluster of all individuals
- Hierarchical Clustering


## Example: Clustering Based on Two Variables

- Example data on height and weight for 9 people.

| Height | Weight |
| :---: | :---: |
| 60 | 84 |
| 62 | 95 |
| 64 | 140 |
| 66 | 155 |
| 68 | 119 |
| 70 | 175 |
| 72 | 145 |
| 74 | 197 |
| 76 | 150 |

## Scatterplot: Plot of Height vs. Weight



Can you find 2 or 3 clusters?

## Scatterplot: Plot of Height vs. Weight



Individuals must be closer within a cluster, but further between clusters. How to measure being close? What type of distance to use?

## Distance Measures for Cluster Analysis

- All clustering methods require the specification of a measure of "similarity". What individuals are considered similar (close) or dissimilar (far)?
- A distance measure is introduced to indicate distances between individuals, and subsequently between clusters
- Some common used distances are:
- Euclidian or Square Euclidian
- Mahalanobis
- Maximum
- Manhattan


## Distance: How "Far" (Dissimilar) is X from Y

- For two subjects X and Y with data $x=\left(x_{1}, x_{2}, \ldots, x_{p}\right)$ and $\mathrm{y}=$ $\left(y_{1}, y_{2}, \ldots, y_{p}\right)$ the following distances can be used to measure the degree of similarity or dissimilarity:
- Euclidian distance:

$$
D(X, Y)=\sqrt{\sum_{i}\left(x_{i}-y_{i}\right)^{2}}
$$

- Mahalanobis distance:

$$
D_{M}(X, Y)=\sqrt{(x-y)^{T} \Sigma^{-1}(x-y)}
$$

( $\Sigma$ is the covariance matrix)

## Partitional Clustering: K-means Method

- Step 1: The $\mathbf{K}$-means partitional clustering method starts with a random selection of K subjects for clusters $C_{1}, C_{2}, \ldots, C_{k}$, where k is determined a priori
- An initial cluster "center" is defined as $T_{k}=X_{k}$, for each cluster
- Step 2: Each subject is assigned to one of these clusters, based on the smallest distance from $T_{k}$ ("center")
- x is assigned to $C_{j}$ if $\mathrm{d}\left(\mathrm{x}, T_{j}\right)$ is the smallest
- Step 3: For the new clusters, calculate the new "centers" $\left(T_{k}=\bar{X}_{k}\right)$ as the means of the subjects in each new cluster
- The procedure (step 2 and step 3 ) is repeated until no subjects are re-assigned


## Partitional Clustering: K-means Method

- K-means is non-hierarchical clustering method. It is faster then hierarchical clustering
- It does not require specification of a linkage method (more on this later)
- The number of clusters, $k$, is pre-specified and fixed
- Hierarchical clustering, on the other hand, provides insight into the clustering process and does not required a pre-specified number of clusters


## Hierarchical Clustering

- All hierarchical clustering methods start with each individual defining its own cluster. Then clusters are joined sequentially in a hierarchical way
- How are two clusters joined:
- Calculate the distance, $\mathrm{D}\left(C_{i}, C_{j}\right)$, between every pair of clusters based on a linkage criteria (more later)
- Then join the two "nearest" clusters who have the smallest $\mathrm{D}\left(C_{i}, C_{j}\right)$
- Continue until there is only one cluster


## Example: Hierarchical Clustering for Height/Weight Data



In 2D and with small data this is possible. With more than 2 variables Dendrogram, is a better way

## Dendograms

- Dendrogram is a useful graphical tool for displaying multidimensional hierarchical structure of clustering
- It shows the distances between individuals (and clusters) in a tree-like structure
- Individuals (or clusters of individuals) that are closest to each other are connected by a horizontal line, forming a new cluster


## Dendrogram Example for Height Weight Data



## Dendrogram Example for Height Weight Data

Merging clusters is based on a linkage criteria:

- Single linkage
- Complete linkage
- Average linkage
- Ward linkage


The distance of a particular pair of objects (or clusters) is reflected in the height of the horizontal line. It is based on the linkage criteria

## Single Linkage Clustering (Minimum)

In single linkage, the distance between two clusters is computed as the distance between the two closest elements in the two clusters:

$$
D(C 1, C 2)=\min _{x \in C 1 ; y \in C 2}\{d(x, y)\}
$$


$D(C 1, C 2)$ is the shortest distance among C1, C2, and C3, so link C1 and C2

## Complete Linkage Clustering (Maximum)

In complete linkage, the distance between two clusters is computed as the distance between the two farthest elements in the two clusters:

$$
D(C 1, C 2)=\max _{x \in C 1 ; y \in C 2}\{d(x, y)\}
$$


$D(C 1, C 3)$ is the shortest distance among C1, C2, and C3, so link C1 and C3

## Average Linkage Clustering (Mean)

In average linkage, the distance between two clusters is computed as the mean of all distances between pairs of elements in the two clusters

$$
D(C 1, C 2)=\operatorname{mean}_{x \in C 1, y \in C 2}\{d(x, y)\}
$$


$D(C 1, C 3)$ is the shortest distance among C1, C2, and C3, so link C1 and C3

## Ward Linkage Clustering

Ward's criterion minimizes the total within-cluster variance. At each step the pair of clusters that result in a minimum increase in variance are merged


# Dendrogram Example for Height Weight Data Using Different Linkage Rule 



Single: Minimum


Average: Mean


Complete: Maximum


Ward


## Illustrating Cluster Analysis Using TROPHY Data

- We will use TROPHY data to group subjects into clusters based on their metabolic risk measures
- Insulin, Glucose, Ins:Gluc ratio, Triglycerides, HDL, LDL, HDL:LDL ratio, Total Cholesterol, Systolic blood pressure, and diastolic blood pressure
- Earlier in PCA we showed that PC1 and PC2 contain most of the information on the metabolic risk
- So, it will be simpler to run cluster analysis based on PC1 and PC2 alone, without losing much of the information of the original data


## Dendrogram for TROPHY Example



## Naming Clusters

- Becouse cluster analysis is an unsupervised process for identifying clusters, giving a name to each cluster it's not easy
- Usually, you look for specific features (based on X1,X2,...,Xp measures) to give an appropriate name
- This may be hard, if clustering is based on a large number of variables
- The data reduction via PCA make it easier, as its focused on features specific to few PC's (i.e. PC1, PC2)






## Heatmap

- A heatmap is a graphical representation of multidimensional structure data using colors
- In a heatmap individual values are distinguished by colors. E.g. large values are colored red, low values yellow
- Dendogram is often added to a heatmap by permuting the rows/columns of a matrix to place similar values near each other
- Examples: DNA microarrays data. Represent the level of expression of many genes across a number of comparable samples



## Summary Points

- Adjustment in Multiple Testing (e.g. testing $m$ hypothesis)
- Family-Wise Error Rate (FWER)
- Bonferroni adjustment (works for small m)
- Set the significance of each test at $p_{k} \leq \frac{\alpha}{m}$. Then FWER $\leq m\left(\frac{\alpha}{m}\right)=\alpha$
- False Discovery Rate (FDR)
- Control the FDR $\leq \alpha$ ( $\alpha$ is an excepted rate of false discoveries)
- For each of $m$ tests get the p -value. Order them: $p_{1} \leq p_{2} \leq \ldots \leq p_{m}$. Find largest k , such that $p_{k} \leq \frac{k * .05}{m}$. Then $p_{1}, p_{2}, \ldots, p_{k}$ are consider significant (reject $H_{01}, \ldots, H_{0 k}$ )


## Summary Points

- Principal component analysis (PCA) is a dimension-reduction technique that looks for structure among variables ( $X_{1}, X_{2}, \ldots, X_{p}$ )
- PCA finds a small number of uncorrelated linear combinations of the variables ( X 's), which summarize most of the information from X's
- Number of PC's. Select the number of PC's based on:
- Eigenvalue criteria: Chose all PC's for which Eigenvalue > 1
- Visually: Use the screen plot to identify the number of PC's where the plot start to flatten
- Explained Variance: Chose a \# of PC's that explain a "sufficient" amount of the variance
- Interpretation of PC's. Use biplots to see how each of the original data (X's) contributes to a PC


## Summary Points

- Cluster Analysis identifies clusters of individuals/objects in a dataset that are similar based on a distance (e.g. Euclidian, Mahalanobis)
- Partition (non-hierarchical method)
- Use K-mean method to find k (pre-specified) clusters
- Hierarchical clustering
- Identify clusters staring with each individual as its own cluster. Next merge clusters (using a linkage criteria) hierarchically until all are part of one cluster
- Common linkage criteria:

Single linkage, complete linkage, average linkage, Ward linkage

- Dendrograms: A visual tree-like structure describing the hierarchical nature of clustering in the data
- Heatmap: A graphical representation of multidimensional structure data using colors

