Descriptions

Models

SAMPLING

VISUAL TESTS

Introduction to Biostatistics



Arranged by Linda Strausbaugh (Genetics 147:5, 1997)

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Contents of Lecture 1-2

Data and descriptive statistics

 $Probability\ theory\ and\ models$

Sampling, SE(M) and CLT.

Visual tests (of normality)

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Introduction to Biostatistics Lecture 1B and 2

Henrik Renlund

January 23-24, 2017



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What shall we learn today (and tomorrow)?

- Data description
 - Graphs
 - Tables and summary measures
- Probability Models
 - Glimpse at theory
 - Normal distribution and some properties
 - Some properties of samples and the Central Limit Theorem.

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Types of data

A data set contains one or more variables, typically arranged as:

Individual Gend		niibamiii	Diabetes	парртнезз
1 M	29	3.92	0	\smile
2 F	37	4.12	0	
3 F	25	4.75	1	_
: :	:	:	:	

Theoretical data categories:

- Numerical
 - discrete; typically integer valued 1, 2, ..., like Age, or
 - continuous; i.e. any value in an interval, like Albumin.
- Categorical
 - nominal, e.g. Gender, or
 - ordinal, e.g. Happiness: $\frown, -, \smile$.

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"Table 1"

It is useful to provide a summary table of the variables you are working with. Choice of descriptive measures may be context dependent.

variable	Diabete	s No	Diabete	s Yes
value	mean	sd	mean	sd
Age	32.0	15.9	32.5	14.1
Albumin	4.20	0.37	3.80	0.50
	percent	n	percent	n
Gender				
M	64%	27	52%	22
F	36%	15	48%	20
Happiness				
	61%	19	36%	15
_	23%	7	36%	15
$\overline{}$	16%	5	36%	12

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How statisticians spend their time

In practice data might be stored in the wrong format.

Check this prior to analysis! This is especially important if data has been transferred (between formats, different OS, etc).

Common problems:

- date- and categorical data suddenly stored as integers
- numerical values stored as text (due to ',' vs. '.')

Usually one only needs to distinguish

- measurements (Age, Albumin are numerical), and
- categories (Gender, Diabetes, Happiness are categorical).

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Some points on tables (for publication)

- Table and caption should be self contained.
- Every table should be referred to in text.*
- Put captions above the table.
- Avoid excessive precision. (Mean age 65.63?)
- Use adequate measures of location and spread.

Generally bad: "Table 3 displays values of biomarkers in group A and B".

Better: "The mean level of measured biomarkers are higher in group A compared to B (Table 3)."

^{*}The table should illustrate some part of the data relating to the hypothesis of the paper. Referring to a table should fit the narrative.

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What about missing data?

Missing data is usually a problem for observational studies.

Typically data is missing in explanatory variables.

Suppose we investigate death within one year of myocardial infarction with a model that includes gender, age, BMI (some missing) and smoking status (some missing).

The statistical software default is to include only those individuals with complete case data on all variables that you include.

Complete case analysis will only give an unbiased result if the reason that a variable is missing has nothing to do with the actual value (and/or the outcome).

If the missing mechanism is known this might be utilized in the analysis.

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Visualization of (continuous) data

A sufficiently small data set might not need visualization.

The level of the protein albumin was recorded in a sample (of size 8) of mice (56 days old):

1.88 2.03 2.11 1.77 2.04 2.05 1.94 1.95

(measured in g/dL - this data set will return in lecture 3).

One simple way to get some handle on data is to order it:

1.77 1.88 1.94 1.95 2.03 2.04 2.05 2.11

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Solutions...ish

There is no trick that guarantees a non-biased analysis.

Single imputation e.g. replace missing value with a "typical" value for that variable (e.g. mean or median). This method underestimates the variance in the variable and will give overly optimistic results.

Multiple imputation create multiple imputed data sets where the missing values are replaced differently in each iteration (e.g. drawn at random from the non-missing values).

"It is not that multiple imputation is so good; it is really that other methods for adressing missing data are so bad."

(Donald Rubin)

N.B. we typically do not impute our outcome data!

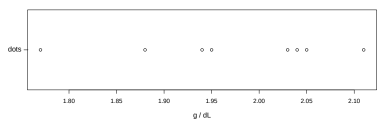
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Dotplot of albumin data

A dotplot is a one dimensional plot of the data.



If there are non-unique (or close) points, the data set may appear smaller than it really is.

This can be alleviated by

- perturbation, or,
- alpha transparency.

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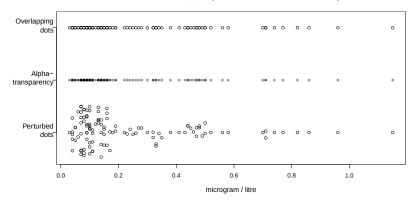
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Subarachnoidal bleeding

A biomarker - the protein $S100\beta$ - was measured for 113 individuals with aneurysmal subarachnoid hemorrhage. (To return in lecture 10.)



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Percentiles

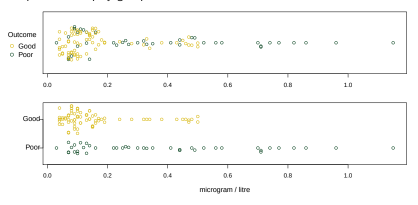
- The kth percentile is a value v such that k percent of your data lies below (or at) v. (Usually not uniquely defined.)
- The 50th percentile (the *median*) is the point which divides your ordered sample equally. (Only 'unique' if sample is odd, else use mean of the two midpoints.)
- The Quartiles: Q1 is the 25th percentile, Q2 is the 50th percentile and Q3 is the 75th percentile.
- We can describe all percentiles with the *empirical cumulative* distribution function (ecdf). When the sample size is 113 the jumps in the ecdf will be multiples of $1/113 \approx 0.01$.

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Dotplot and groups

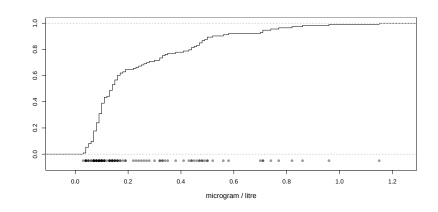
Dotplots can display groups.



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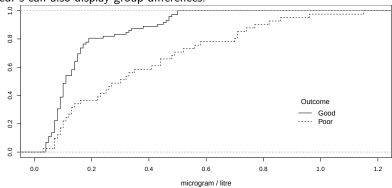
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Empirical cumulative distribution function for $S100\beta$



Empirical cumulative distribution function

Ecdf's can also display group differences.



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Mean or median?

Ex: A small company has 5 employees, who earns 19, 21, 22, 24, 27 (K SEK) and a boss who earns 55.

Salaries	Employees	Entire company
Median	22	23
Mean	22.6	28

Some points:

- Small data sets might not need summary measures.
- Symmetric data has mean \approx median.
- (Easy enough to calculate both.)

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Average value / Measure of location

An average value is a single value meant to be representative of the entire data set.

- **The mode:** is the single most common value of a data point. (Mostly meaningful for categorical data.)
- The median: is the midpoint of the ordered numerical sample.
- The mean: is the "center of gravity" of a data set.
 Note: unlike the median, the mean value is sensitive to "extreme" values.

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Measuring the spread of a data set

- Range The difference between the maximum and the minimum value.
- Interquartile range (IQR): Q3-Q1.
- Standard deviation (sd) is given by the formula,

$$s = \sqrt{\frac{1}{n-1} \sum_{i=1}^{n} (x_i - \bar{x})^2}.$$

Where x_1, x_2, \dots, x_n is the sample and

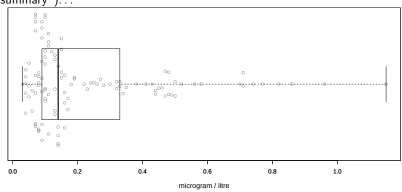
$$\bar{x} = \frac{1}{n} \sum_{i=1}^{n} x_i$$

is the (sample) mean.

It is (approximately) the mean distance to the mean value.

Boxplot of $S100\beta$

The boxplot usually show min, Q1, med, Q3 and max (the "5-point summary")...



Connection between boxplot and ecdf

The boxplot contains less information.

8

9

0.0

0.2

0.4

0.6

0.8

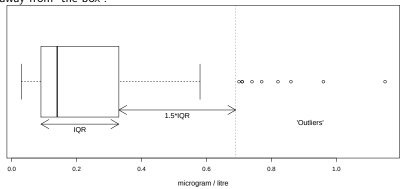
1.0

12

microgram / litre

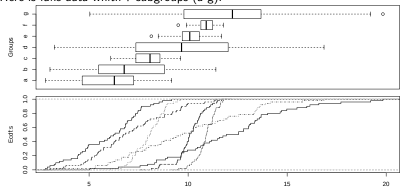
Boxplot

 \dots but most software mark points that are more than 1.5 times the IQR away from 'the box'.



Pattern or detail?

Here is fake data whith 7 subgroups (a-g).



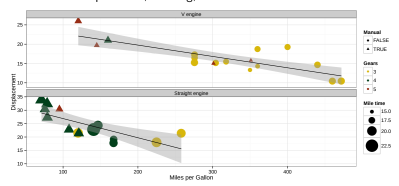
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Advanced graphics...

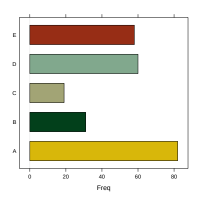
Anyone seriously interested in graphics might want to look at Wilkinson's *The Grammar of Graphics*, which provides a framework for thinking about data visualization. These ideas are implemented in the R package **ggplot2** - which provides tools for mapping variables to aestic properties, adding layers of statistical computations, faceting, etc.

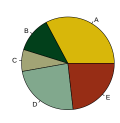


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In defense of pie charts?

Is it of interest to be able to compare summation of levels? (Is A + B + C smaller than D + E?)





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Some points on graphs (for publication)

- Graph and caption should be self-contained.
- Every graph should be referred to in the text.
- 'Economy' Do not make a graph which is more easily expressed in text or a small table.

E.g. graph with a single boxplot.

• Pattern or detail?

E.g.

- a ecdf gives a lot of detail of a data set.
- graph with multiple boxplots can reveal pattern among subgroups.
- Avoid 2D graphs shown in 3D.
- Avoid pie charts? It depends...

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Probability theory studies models of random data. A **model** is a way of specifying the range of possible values and the probability with which these occur.

- Probability functions describe discrete numeric/categorical data
- Density functions describe continuous (numeric) data

The **cumulative distribution function** is a universal way of describing all random (numeric) data. (Will not be discussed.)

- Probability theory: given model (model parameters, or other aspects)
 describe how data behave. E.g.
 - specific results: how likely are deviations from what we expect
 - general results: Law of Large Numbers, Central Limit Theorem, etc.
- **Inference theory:** given data what is a likely model/parameters or other aspects of the underlying distribution (without specifying model = non-parametric statistics).

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Probability models for categorical or integer-valued data

A yet undetermined random value is called a random variable (RV).

Let Z ='the outcome of the throw of a die'. Then $\mathbf{Prob}(Z=k)=1/6$ for all $k=1,2,\ldots,6$, or, equivalently

Value k	1	2	3	4	5	6
Prob(Z = k)	1/6	1/6	1/6	1/6	1/6	1/6

Suppose that in the population there are 49 % non-smokers, 20 % former smokers and 31% current smokers. Then the smoking status X of a person selected at random is a RV with a probability function

Value v	non	former	current
Prob(X = v)	0.49	0.20	0.31

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FEV data set

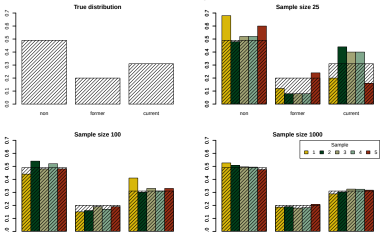
430 children (9-17 years of age) had their age, forced expiratory volume in 1 second (FEV) and smoking status (!) recorded. (To be seen again in lecture 8.)

A barchart is a way to visualize a variable with a small number of unique values (often categorical). They are visual analogous of tables.

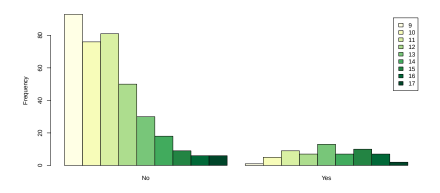
Ex: how do the ages distribute over smoking status?

Smoking	Age								
	9	10	11	12	13	14	15	16	17
No	93	76	81	50	30	18	9	6	6
Yes	1	5	9	7	13	7	10	7	2

Five samples from three different sampling sizes from previous distribution



Visualizing 'age' versus 'smoking'



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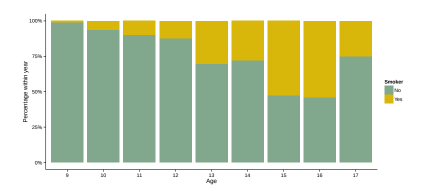
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If the groups (smokers/non-smokers) aren't balanced it is difficult to compare the distributions.

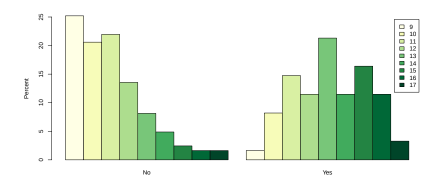
Tabulate/plot the percentages within groups:

Age	Smokii	ıg (%)				
	No	Yes				
9	0.252	0.016				
10	0.206	0.082				
11	0.220	0.148				
12	0.136	0.115				
13	0.081	0.213				
14	0.049	0.115				
15	0.024	0.164				
16	0.016	0.115				
17	0.016	0.033				
Sum	1.000	1.000				

Visualizing 'smoking' versus 'age' as percentages within age groups, stacked



Visualizing 'age' versus 'smoking' as percentages within smoking groups



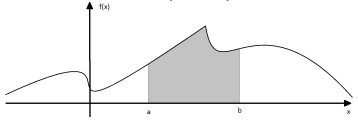
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$Probability\ model\ for\ continuous\ data$

Recall that a RV is a yet undetermined value among several possible numbers.

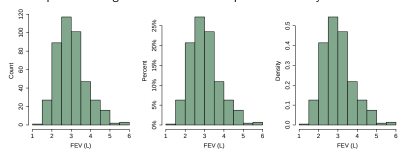
A continuous RV is described by its *density function*.



If X has density function f as above, then we compute probabilities as

Prob
$$(a \le X \le b) = \text{Area}(a,b).$$

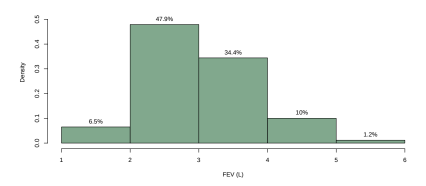
The shape of a histogram estimates the shape of the density function.



- "Density" is more abstract but
 - gives right scale for density function estimate (easy to correctly plot candidate model on top of histogram)
 - allows for varying "bins"
 - allows for comparison between very different sample sizes

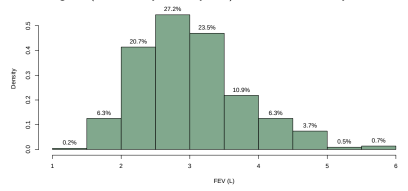
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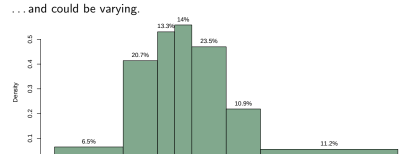
The interval width is arbitrary...



Histogram (with density) of FEV

The histogram (with density on the y-axis) estimates the density function.





FEV (L)

0.0

Sampling

Visual tests

Summary

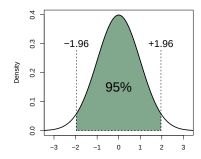
Graph	Summary Measure	Theory
Ecdf	percentiles	(Cdf)
Boxplot	min, Quartiles, max	
Bar charts		Probability functions (discrete
		RV)
Histograms		Density functions (continuous
		RV)
	median, IQR	any distribution
	mean, s.d.	symmetrical distribution ($pprox$
		Normal distribution)

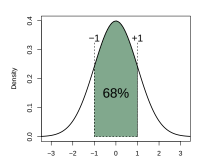
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Properties of the standard Normal distribution (and why you need to know the number 1.96)

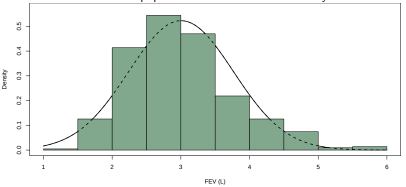
The standard Normal distribution has a standard deviation of 1. 68% of the 'probability mass' lies within ± 1 . 95% of the 'probability mass' lies within ± 1.96 .





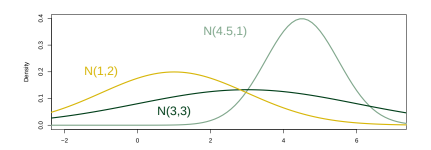
Assuming Normality

Often we assume that the population follows a Normal density curve.



The Normal distribution $N(\mu, \sigma)$

is determined by its mean (μ) and standard deviation (σ) . If X is N(0,1) then $\mu + \sigma X$ is N(μ , σ).



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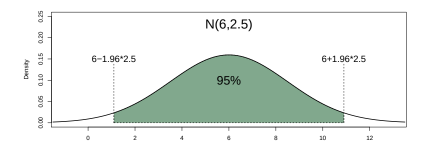
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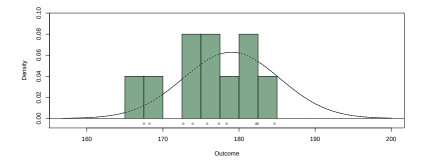
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Properties of the Normal distribution

If X is N(μ , σ), then 95% of observations will be between $\mu-1.96\sigma$ and $\mu+1.96\sigma$.



n = 10

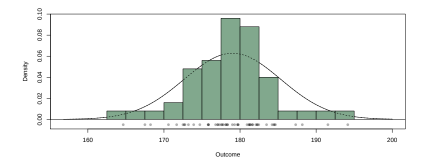


Attempt to visualize sampling from a given model

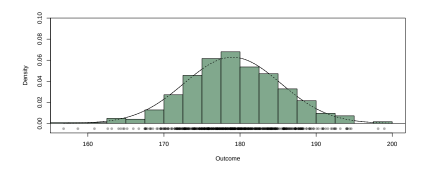
Assume that height of individuals in some population

- is Normally distributed,
- has mean (μ) 179 (cm), and,
- has s.d. (σ) 6.35 (cm).

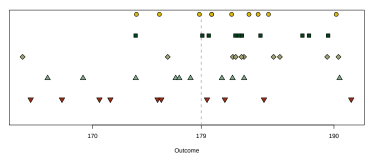
The following three slides show histogram of samples of sizes 10, 50 and 500, respectively.



$$n = 500$$



5 samples of size 10 with population mean = 179 and sd. = 6.35



Properties of samples and sample means

Suppose we have a sample of size 10 from the population.

We want to know the population mean (μ) .

We can estimate with the sample mean(μ^*).

But how good a guess is μ^* ?

Well...if the sample is drawn at random¹ then μ^* is *unbiased* (on average correct).

If we knew that the sd of $\mu^*=180$ was, say 3.0, then there is an (approximate!) interpretation.

The estimated value 180 (cm) is on average off by 3.0 (cm).

But what *is* the standard deviation of a sample mean?

Consider several iterations of the procedure of drawing a sample of size 10.

Properties of sample means

We can see that samples and their means and standard deviations vary (of course).

Standard Error of the Mean (SEM)

The standard deviation of the sample mean is referred to as SEM and a measure of (approx.) the average distance the population mean. Thus a measure of 'how good' the sample mean is as an estimator of the population mean.

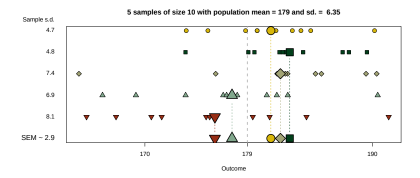
In a Normal model the SEM is the population s.d. dived by \sqrt{n} .

The next slide tries to visualize this.

¹All possible samples are equiprobable

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$Normal\ population \Rightarrow Normal\ mean$

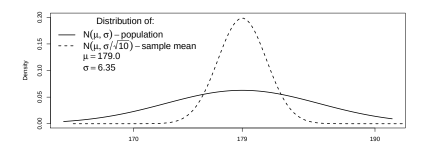
On the previous slide you saw

- a Normal population curve, and
- the Normal curve for the sample mean (for sample size 10).

In this case this is the true (derived from theory) density curve for the sample mean.

Properties of the Normal distribution will aid further analysis using the mean of the sample.

What happens if the population follows some non-Normal curve?



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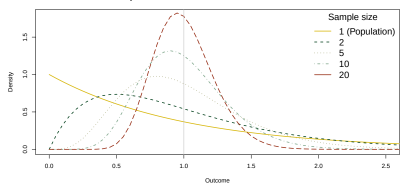
The Central Limit Theorem (CLT)

CLT: Regardless of the true population density curve, the sample mean density can be made (with arbitrarily good approximation) Normal by choosing n large enough.

- How large does n have to be?
 Depends on how skew the population density is.
 - In general n = 20 will suffice.
 - If population is Normal then n = 1 is enough.
- CLT applies to many 'statistics' (= functions of samples).

Attempt to visualize CLT for the sample mean

The distribution of sample means from a skewed distribution.



Cross-over data (from lecture 4)

13 patients had their peak expiratory flow (PEF, I/min) recorded once after inhaling each of two different asthma drugs (the order of which were random).

In *paired* data one usually look at the 13 differences as a measurement of differences in individual effect size.

Data:

40, 50, 70, 20, 40, 30, -35, 15, 90, 30, 30, 80, 130

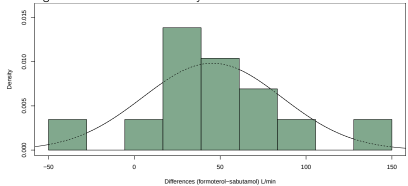
Is the normal distribution a good model for these 13 numbers?

Quantiles?

Quantiles divides your data into (roughly) equal piles.

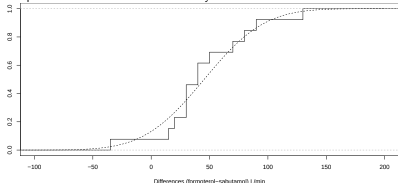
- the median is the 2-quantile
- the tertiles are the 3-quantiles (the $33\frac{1}{3}$ percentile and the $66\frac{2}{3}$ percentile)
- the quartiles (Q1, Q2 and Q3) are the 4-quantiles.
- ...and so on.

A histogram with best normal density fit.

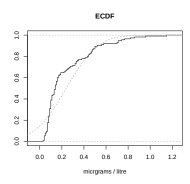


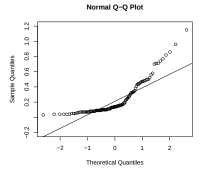
PEV

A empirical cdf with best normal density cdf.



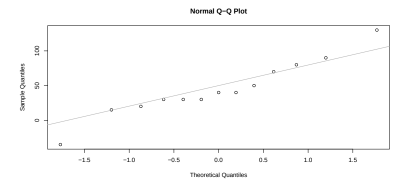
These visual test require some training. There are also formal tests of normality (e.g. Shapiro-Wilks) The S100 β measurements from the subarachnoidal bleeding example is certainly not normally distributed.





The Quantile-Quantile plot

If the differences in effect size is sampled from a Normally distributed population its QQ-plot should be a straight line (approximately). A QQ-plot plots the sample of size *n* against the (slightly shifted) *n*-quantiles of the (standard) Normal distribution.



A reminder ahead of time

It is very rarely the actual data that is tested for normality!

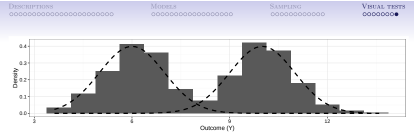
Most of the time the models that assume normality does so for the "error" terms, i.e. there is a model, depending on the covariates x, for the outcome Y such that

Y = some deterministic function of x + noise.

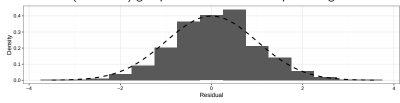
E.g. a 2-sample *t*-test assumes that an outcome is normally distributed around a group-specific mean. Data for such a test might look like this

outcome (Y)					4.7	
group (x)	Α	Α	В	В	Α	

We cannot test the entire Y data for normality. This is evident if imagine the group effect to be very large...



It is the deviations (noise/error term) around each group-specific mean that is supposed to be normal, an estimation of which is called *residuals*. Subtract the (estimated) group effect from each datapoint to get:



References

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Second Summary

95% of observations from a Normal population lies within 1.96 multiples of the (population) s.d. from the (population) mean.

Means and in particular s.d. must be distinguished on three levels

- population,
- sample, and
- estimate.

The s.d. of the latter is called the Standard Error (Standard Error of the Mean if the mean is used as estimate.)

(The sample s.d. is an estimate of the population s.d.)

The CLT explains why many estimates ('statistics') are (approx.) Normally distributed even though the population may not be.