# BST02: Using R for Statistics in Medical Research

Part D: Statistics with R

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#### t-test: t.test()

#### One-sample t-test

ightharpoonup compares the mean of a sample with a fixed value  $\mu$ 

#### Two sample / independent samples t-test

 $\blacktriangleright$  compares the difference between the means of two samples with a fixed value  $\mu$ 

#### **Related samples t-test**

ightharpoonup compares the mean of the difference between related observations with a fixed value  $\mu$  (same as one-sample t-test)

## Wilcoxon Test: wilcox.test()

#### **Wilcoxon Signed Rank Test**

lacktriangle tests if one sample (or the differences between two paired samples) is/are symmetric about  $\mu$ 

#### Wilcoxon Rank Sum Test / Mann-Whitney test

► test for a location shift between the distributions of two independent samples

See also BBR Sections 7.2 & 7.3 (http://hbiostat.org/doc/bbr.pdf)

### Kruskal-Wallis Rank Sum Test: kruskal.test()

- ► This is an extension of the Wilcoxon rank sum test for more than two groups
- Test for a difference in location of a continuous variable between multiple groups
- ► The Wilcoxon rank sum test is a special case of the Kruskal-Wallis rank sum test

#### Other tests for continuous data

- ► Kolmogoriv-Smirnov Test: ks.test() tests if two samples are drawn from the same continuous distribution
- ► Shapiro-Wilk Normality Test: shapiro.test()
- ► Friedman Rank Sum Test: friedman.test() non-parametric test for 2 or more related samples
- **.**..

# **Tests for Categorical Data / Proportions**

#### **One-sample Proportion Test**

- tests if the proportion in one sample is equal to a fixed value p
- prop.test() and binom.test()

#### **Tests for Proportions in Multiple (independent) Groups**

- tests if the proportion in several samples are equal
- chisq.test() and fisher.test() (when there are cells with 0)

See also BBR Sections 5.7 & 6 (http://hbiostat.org/doc/bbr.pdf)

# **Tests for Categorical Data / Proportions**

#### **Related Samples: McNemar Test**

- ▶ Tests for symmetry in a  $2 \times 2$  table
- mcnemar.test()

#### **3-Dimensional Contingency Table**

- Cochrane-Mantel-Haenszel Test
- $\triangleright$   $\chi^2$  test for independence of two nominal variables in each stratum
- ▶ mantelhaen.test()

#### **Statistical Tests**

#### **Demos**

- ▶ Name of the demo
- ▶ Name of demo

#### **Practicals**

- ► Name of practical html
- ► Name of practical html

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#### **Statistical Tests**

#### Continuous Outcomes

- ▶ t.test()
- ▶ wilcox.test()
- kruskal.test()
- ks.test()
- ▶ friedman.test()
- shapiro.test()

#### **Categorical Outcomes**

- prop.test()
- ▶ binom.test()
- chisq.test()
- ▶ fisher.test()
- mcnemar.test()
- mantelhaen.test()

# Variance and Correlation

- cor.test()
- ▶ bartlett.test()
- var.test()

#### **Pairwise tests**

- pairwise.prop.test()
- pairwise.t.test()
- pairwise.wilcox.test()

# **Linear Regression**

A standard linear regression model has the form

$$y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \dots + \beta_p x_p + \varepsilon$$
 with  $\varepsilon \sim N(0, \sigma^2)$ 

#### where

- ▶ *y* is the **outcome** variable ("dependent variable")
- $\triangleright$   $x_1, \dots, x_p$  are the **covariates** ("independent variables")
- $ightharpoonup \beta_0, \dots, \beta_p$  are the **regression coefficients** 
  - $\triangleright$   $\beta_0$  is the intercept
  - $ightharpoonup \beta_1, \dots, \beta_p$  estimate the effects of the covariates
- ightharpoonup arepsilon is a vector of **residuals**, which we assume to be (approximately) normally distributed.

# **Linear Regression**

To fit a **linear regression** in R we use the function lm().

The most important arguments are

- formula:
   a formula object
- data: a data.frame (optional, but usually needed)
- subset: a vector specifying which observations should be used (optional)

#### **Model Formula**

A formula object has the form

outcome ~ linear predictor

for example

$$y \sim x1 + x2 + x3$$

- ► Variables are separated by "+" signs.
- ► An intercept is automatically included.
- one-sided formulas (omitting the outcome) are possible (used for random effects specification)

#### **Model Formula: Interactions**

Interaction terms are written using ":" or "\*".

"\*" includes the main effects and interaction terms, i.e.,

is equivalent to

$$y \sim x1 + x2 + x1:x2$$

Interactions between multiple variables can be written using "()", i.e.,

$$y \sim x1 * (x2 + x3)$$

is equivalent to

$$y \sim x1 * x2 + x1 * x3$$

#### **Model Formula: Interactions**

To specify a **higher level interaction** (for example a three-way interaction) "^" is used, i.e.,

$$y \sim (x1 + x2 + x3)^3$$

will create all interactions up to 3-way and is equivalent to

$$y \sim x1 * x2 * x3$$

and equivalent to

$$y \sim x1 + x2 + x3 + x1:x2 + x1:x3 + x2:x3 + x1:x2:x3$$

and

$$v \sim (x1 + x2 + x3)^2$$

will create all two-way interactions and is equivalent to

$$y \sim x1 + x2 + x3 + x1:x2 + x1:x3 + x2:x3$$

# **Model Formula: Removing terms**

The "-" sign can be used to remove terms from a model formula, for example

is equivalent to

$$y \sim x1 + x3 + x1:x2 + x2:x3 + x1:x2:x3$$

The **intercept** can be removed from a formula by using "-1" or "+0", i.e.

$$y \sim x1 + x2 - 1$$
  
 $y \sim x1 + x2 + 0$ 

# **Generalized Linear Regression (GLM)**

A generalized linear regression model has the form

$$g(\mathbb{E}(y)) = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \dots + \beta_p x_p$$

where g() is a link function and y is from the exponential family.

For example **logistic regression** for binary *y*:

$$log\left(\frac{P(y=1)}{1-P(y=1)}\right) = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \dots + \beta_p x_p$$

 $log\left(\frac{p}{1-p}\right)$  is the **logit** link.

# **Generalized Linear Regression (GLM)**

To fit a GLM in R we use the function glm().

The most important arguments are

- formula: a formula object
- family: a family object or name of the family function, describing the error distribution and link function
- data: a data.frame (optional, but usually needed)
- subset: a vector specifying which observations should be used (optional)

#### **Families and Link Functions**

Common families & available links in R:

(see also ?family)

| family   | link                                 |
|----------|--------------------------------------|
| binomial | logit, probit, cauchit, log, cloglog |
| gaussian | identity, log, inverse               |
| Gamma    | inverse, identity, log               |
| poisson  | log, identity, sqrt                  |

The family argument in glm() can be specified in the following ways:

- binomial(link = "logit")
- binomial()
- ▶ binomial
- ▶ "binomial"

#### Note:

When the link is not explicitely specified (i.e. option 1), the default link is used.

# **Regression Basics**

#### **Demo**

► Regression Basics R html

#### **Practical**

- ► Linear Regression html
- ► Name of practical html

# Regression

#### **Regression Models**

- ► lm()
- **▶** glm()

#### **Regression Results**

- summary()
- coef(), confint()
- fitted(), residuals()
- ► AIC(), BIC()
- anova()

#### **Plots**

- ▶ plot()
- qqnorm(), qqline(), qqplot()

#### **Topic**

- ▶ ns(), bs(), I()
- p.adjust()
- ▶ all.vars()
- update()
- ▶ as.formula()