## REGRESSION MODELS

## ANOVA MODELS




Uses dummy variables to represent categorical variables!

## Outline

- Motivation
- ANOVA as a regression model
- Dummy variables
- One-way ANOVA models
- Contrasts
- Multiple comparisons
- Two-way ANOVA models
- Interactions
- ANCOVA models
- Experimental Designs and ANOVA models


## Motivation

## Motivation

- Let's investigate if genetic factors are associated with cholesterol levels.
- Ideally, you would have a confirmatory analysis of scientific hypotheses formulated prior to data collection
- Alternatively, you could consider an exploratory analysis - hypotheses generation for future studies


## ANOVA/ANCOVA: Motivation

- Scientific hypotheses of interest:
- Assess the effect of rs174548 on cholesterol levels.
- Assess the effect of rs174548 and gender on cholesterol levels
- Does the effect of rs174548 on cholesterol differ between males and females?
- Assess the effect of rs174548 and age on cholesterol levels
- Does the effect of rs174548 on cholesterol differ depending on subject's age?
- Scientific question:
- Assess the effect of rs174548 on cholesterol levels.


## Motivation: Example

Here are some descriptive summaries:


## Motivation: Example

Another way of getting the same results:

```
> by(chol, as.factor(rs174548), mean)
as.factor(rs174548): 0
[1] 181.0617
as.factor(rs174548): 1
[1] 187.8639
as.factor(rs174548): 2
[1] 186.5
> by(chol, as.factor(rs174548), sd)
as.factor(rs174548): 0
[1] 21.13998
as.factor(rs174548): 1
[1] 23.74541
-----------------------------------------------------------------------------
as.factor(rs174548): 2
[1] 17.38333
```



## Motivation: Example

Another graphical display:


## Motivation: Example

- Feature:
- How do the mean responses compare across different groups?
- Categorical/qualitative predictor

ANOVA

As a regression model

## ANalysis Of VAriance Models (ANOVA)

- Compares the means of several populations


Independence
Normality
Equal variances

## ANalysis Of VAriance Models (ANOVA)

- Compares the means of several populations



## ANalysis Of VAriance Models (ANOVA)

- Compares the means of several populations
- Counter-intuitive name!


## ANalysis Of VAriance Models (ANOVA)

In both data sets, the true population means are: 3 (A), 5 (B), 7(C)


Where do you expect to detect difference between population means?

## ANalysis Of VAriance Models (ANOVA)

- Compares the means of several populations
- Counter-intuitive name!
- Underlying concept:
- To assess whether the population means are equal, compares:
- Variation between the sample means (MSR) to
- Natural variation of the observations within the samples (MSE).
- The larger the MSR compared to MSE the more support that there is a difference in the population means!
- The ratio MSR/MSE is the F-statistic.


## Decomposition of sum of squares



## ANalysis Of VAriance Models (ANOVA)

- Equivalent to regression with categorical predictors.
- Predictors represented with "dummy" variables


## ANOVA as a multiple regression model

- Dummy Variables:
- Suppose you have a categorical variable C with k categories. To represent that variable we can construct k -1 dummy variables of the form
$x_{1}=\left\{\begin{array}{l}1, \text { if subject is in category } 2 \\ 0, \text { otherwise }\end{array}\right.$
$x_{2}=\left\{\begin{array}{l}1, \text { if subject is in category } 3 \\ 0, \text { otherwise }\end{array}\right.$
$x_{k-1}=\left\{\begin{array}{l}1, \text { if subject is in category k } \\ 0, \text { otherwise }\end{array}\right.$
The omitted category (here category 1 ) is the reference group.


## ANOVA as a multiple regression model

- Dummy Variables:
- Back to our motivating example:
- Predictor: rs174548 (coded $0=\mathrm{C} / \mathrm{C}, 1=\mathrm{C} / \mathrm{G}, 2=\mathrm{G} / \mathrm{G}$ )
- Outcome (Y): cholesterol

Let's stake C/C as the reference group.

$$
\begin{aligned}
& x_{1}= \begin{cases}1, & \text { if code } 1(\mathrm{C} / \mathrm{G}) \\
0, \text { otherwise }\end{cases} \\
& x_{2}= \begin{cases}1, & \text { if code } 2(\mathrm{G} / \mathrm{G}) \\
0, \text { otherwise }\end{cases}
\end{aligned}
$$

ANOVA as a multiple regression model

| rs174548 | $X_{1}$ | $X_{2}$ |
| :--- | :--- | :--- |
| C/C | 0 | 0 |
| C/G | 1 | 0 |
| G/G | 0 | 1 |

## ANOVA as a multiple regression model

- Regression with Dummy Variables:
- Example:

Model: $E\left[Y \mid x_{1}, x_{2}\right]=\beta_{0}+\beta_{1} x_{1}+\beta_{2} x_{2}$

- Interpretation of model parameters?

ANOVA as a multiple regression model

- Regression with Dummy Variables:
- Example:

Model: $\mathrm{E}\left[\mathrm{Y} \mid \mathrm{x}_{1}, \mathrm{x}_{2}\right]=\beta_{0}+\beta_{1} \mathrm{x}_{1}+\beta_{2} \mathrm{x}_{2}$

- Interpretation of model parameters?
- $\beta_{0}$ : mean cholesterol when rs174548 is C/C
- $\beta_{0}+\beta_{1}$ : mean cholesterol when rs174548 is C/G
- $\beta_{0}+\beta_{2}$ : mean cholesterol when rs174548 is G/G


## ANOVA as a multiple regression model

- Regression with Dummy Variables:
- Example:

Model: $\mathrm{E}\left[\mathrm{Y} \mid \mathrm{x}_{1}, \mathrm{x}_{2}\right]=\beta_{0}+\beta_{1} \mathrm{X}_{1}+\beta_{2} \mathrm{X}_{2}$

- Interpretation of model parameters?
- $\beta_{0}$ : mean cholesterol when rs174548 is C/C
- $\beta_{0}+\beta_{1}$ : mean cholesterol when rs174548 is C/G
- $\beta_{0}+\beta_{2}$ : mean cholesterol when rs174548 is G/G
- Alternatively
- $\beta_{1}$ : difference in mean cholesterol levels between groups with rs174548 equal to $C / G$ and $C / C$.
- $\beta_{2}$ : difference in mean cholesterol levels between groups with rs174548 equal to $\mathrm{G} / \mathrm{G}$ and $\mathrm{C} / \mathrm{C}$.


## ANOVA as a multiple regression model

- Alternative parameterization
- Each group with its own mean!
- Let' s re-write the model:

$$
\text { Model: } \mathrm{E}\left[\mathrm{Y}_{\mathrm{ij}}\right]=\mu_{\mathrm{i}}
$$

(i: genotype index, $\mathrm{j}:$ subject index)

ANOVA as a multiple regression model

- Regression Model:

Model 1: $E\left[Y \mid x_{1}, x_{2}\right]=\beta_{0}+\beta_{1} x_{1}+\beta_{2} x_{2}$.

- ANOVA Model:

Model 2: $\mathrm{E}\left[\mathrm{Y}_{\mathrm{ij}}\right]=\mu_{\mathrm{i}}$

ANOVA as a multiple regression model

| Mean | Regression <br> Model |
| :---: | :---: |
| $\mu_{1}$ | $\beta_{0}$ |
| $\mu_{2}$ | $\beta_{0}+\beta_{1}$ |
| $\mu_{3}$ | $\beta_{0}+\beta_{2}$ |

## ANOVA as a multiple regression model

- Regression Model:

Model 1: $\mathrm{E}\left[\mathrm{Y} \mid \mathrm{x}_{1}, \mathrm{x}_{2}\right]=\beta_{0}+\beta_{1} \mathrm{x}_{1}+\beta_{2} \mathrm{x}_{2}$.

- ANOVA Model:

Model 2: $\mathrm{E}\left[\mathrm{Y}_{\mathrm{ij}}\right]=\mu_{\mathrm{i}}$

Key Message:
ANOVA is a special case of a regression model!

ANOVA as a multiple regression model

- The same idea applies to problems with several categorical predictors [aka: factors]
- One-way ANOVA: one factor
- Two-way ANOVA: two factors
- ...
- Model assumptions
- Equal variances
- Normality
- Independence


## One-way ANOVA models

## ANOVA: One-Way Model

- Goal:
- Compare the means of K independent groups (defined by a categorical predictor)
- Statistical Hypotheses:
- (Global) Null Hypothesis:
$H_{0}: \mu_{1}=\mu_{2}=\ldots=\mu_{K}$.
- Alternative Hypothesis:
$H_{1}$ : not all means are equal
- If the means of the groups are not all equal (i.e. you rejected the above $\mathrm{H}_{0}$ ), determine which ones are different (multiple comparisons)


## Estimation and Inference

- Global Hypotheses
$\mathrm{H}_{0}: \quad \mu_{1}=\mu_{2}=\ldots=\mu_{K}$
vs. $\quad H_{1}$ : not all means are equal
- Analysis of variance table

| Source | df | SS | MS | $F$ |
| :--- | :--- | :--- | :--- | :--- |
| Regression | $\mathrm{K}-1$ | $\mathrm{SSR}=\sum_{i}\left(\overline{\mathrm{y}}_{\mathrm{i}}-\overline{\mathrm{y}}\right)^{2}$ | MSR= <br> SSR/(K-1) | MSR/ <br> MSE |
| Residual | $\mathrm{n}-\mathrm{K}$ | $\mathrm{SSE}=\sum_{i, j}\left(\mathrm{y}_{\mathrm{ij}}-\overline{\mathrm{y}}_{\mathrm{i}}\right)^{2}$ | MSE $=$ <br> SSE/n-K |  |
| Total | $\mathrm{n}-1$ | $\mathrm{SST}=\sum_{\mathrm{i}, j}\left(\mathrm{y}_{\mathrm{ij}}-\overline{\mathrm{y}}\right)^{2}$ |  |  |

## ANOVA as a multiple regression model

Back to example:

| Mean | Regression Model |
| :---: | :---: |
| $\mu_{1}$ | $\beta_{0}$ |
| $\mu_{2}$ | $\beta_{0}+\beta_{1}$ |
| $\mu_{3}$ | $\beta_{0}+\beta_{2}$ |

## Estimation and Inference

- Global Hypotheses
$\mathrm{H}_{0}: \beta_{1}=\ldots=\beta_{K-1}=0$
vs. $\quad H_{1}$ : not all coeffs are zero
- Analysis of variance table

| Source | df | SS | MS | F |
| :--- | :--- | :--- | :--- | :--- |
| Regression | $\mathrm{K}-1$ | $\mathrm{SSR}=\sum_{i}\left(\overline{\mathrm{y}}_{\mathrm{i}}-\overline{\mathrm{y}}\right)^{2}$ | MSR $=$ <br> SSR/(K-1) | MSR/ <br> MSE |
| Residual | $\mathrm{n}-\mathrm{K}$ | $\mathrm{SSE}=\sum_{i, j}\left(\mathrm{y}_{\mathrm{ij}}-\overline{\mathrm{y}}_{\mathrm{i}}\right)^{2}$ | MSE $=$ <br> SSE/n-K |  |
| Total | $\mathrm{n}-1$ | $\mathrm{SST}=\sum_{i, j}\left(\mathrm{y}_{\mathrm{ij}}-\overline{\mathrm{y}}\right)^{2}$ |  |  |

## ANOVA: One-Way Model

- How to fit a one-way model as a regression problem?
- Need to use "dummy" variables
- Create on your own (can be tedious!)
- Most software packages will do this for you
- R creates dummy variables in the background as long as you state you have a categorical variable (may need to use: as.factor)

| - ANOVA: One-Way Model |  |
| :---: | :---: |
| $\begin{aligned} & \text { By hand: } \\ & \text { Creating "dummy" } \end{aligned}$ variables: | > fit0 = lm(chol ~ dummy1 + dummy2) > summary(fit0) Call: $\operatorname{lm}($ formula = chol ~ dummy1 + dummy2) Residuals: |
| $>$ dummy $1=1^{*}(r s 174548==1)$ <br> $>$ dummy $2=1^{*}(r s 174548==2)$$\quad$ Coefficients: $\quad$ Estimate Std. Error $t$ value $\operatorname{Pr}(>\|t\|)$ |  |
|  | $\qquad$ |
| Fitting the | danyet of varimes mate |
|  |  |



## ANOVA: One-Way Model

- Your turn!
- Compare model fit results (fit0 \& fit1.1) What do you conclude?


| ANOVA: <br> One-Wa | Model |
| :---: | :---: |
| $\begin{aligned} & >\text { fit } 0=1 \mathrm{~m}(\text { chol } \sim \text { dummy } 1+\text { dummy } 2) \\ & >\text { summary }(\text { fit } 0) \end{aligned}$ | > fitl.1 $=\operatorname{lm}$ (chol $\sim$ as.factor(rs174548)) |
| Call: <br> lm(formula $=$ chol $\sim$ dummy1 + dummy2) | Call: <br> $\operatorname{lm}($ formula $=$ chol $\sim$ as.factor (rs174548)) |
| Residuals: <br> Min 12 Median $3 Q$ Max | Residuals: Min 10 Median a |
| $\begin{array}{llllll}-64.06167 & -15.91338 & -0.06167 & 14.93833 & 59.13605\end{array}$ | $\begin{array}{llrrr}-64.06167 & -15.91338 & -0.06167 & 14.93833 & 59.13605\end{array}$ |
| Coefficients: | Coefficients: <br> Estimate Std. Error $t$ value $\operatorname{Pr}(>\|t\|)$ |
|  Estimate Std. Error t value $\operatorname{Pr}(>\|t\|)$  <br> (Intercept)   <br> 181.062 1.455 $124.411<2 e-16 * * *$ |  |
| $\begin{array}{lllll}\text { dummy } 1 & 6.802 & 2.321 & 2.930 & 0.00358\end{array}$ | $\begin{array}{lrrr} \\ \text { as.factor(rs174548) } 1 & 6.802 & 2.321 & 2.930\end{array}$ |
| $\begin{array}{lllll}\text { dummy2 } & 5.438 & 4.540 & 1.198 & 0.23167\end{array}$ | $\begin{array}{llllll}\text { as.factor(rs174548)2 } & 5.438 & 4.540 & 1.198 & 0.23167\end{array}$ |
| --- |  |
| Residual standard error: 21.93 on 397 degrees of freedom Multiple R-squared: 0.0221 , Adjusted R-squared: 0.01718 F-statistic: 4.487 on 2 and 397 DF, p-value: 0.01184 | Residual standard error: 21.93 on 397 degrees of freedom Multiple R-squared: 0.0221, Adjusted R-squared: 0.01718 F-statistic: 4.487 on 2 and 397 DF, p-value: 0.01184 |
| > anova(fit0) <br> Analysis of Variance Table | > anova(fit1.1) |
| Response: chol | Response: chol Df Sum Sq Mean Sq F value $\operatorname{Pr}(>F)$ |
| Df Sum Sq Mean Sq F value Pr (>F) |  |
| dummy1 1 3624 3624 7.5381 0.006315 ** <br> dummy2 1 690 690 1.4350 0.231665  | $\begin{array}{lllllll}\text { as.factor(rs174548) } & 2 & 4314\end{array}$ |
|  | Residuals $397190875 \quad 481$ |
| --- |  |
| ```> 1-pf(4.4865,2,397) [1] 0.01183671 > 1-pf(((3624+690)/2)/481,2,397) [1] 0.01186096``` | 184 |


| ANOMA: | del |
| :---: | :---: |
|  |  |
| ```> fit1.1 = lm(chol ~ as.factor(rs174548)) > summary(fit1.1)``` | Let' s interpret the regression model results! |
| $\operatorname{lm}($ formula $=$ chol $\sim$ as.factor $($ rs174548) ) | - What is the interpretation of the regression model coefficients? |
| Residuals: |  |
| $\begin{array}{llllll}-64.06167 & -15.91338 & -0.06167 & 14.93833 & 59.13605\end{array}$ |  |
| Coefficients: |  |
| $\begin{array}{lrl} & \text { Estimate } \\ \text { (Intercept) } & \text { Std. Error t value } & \operatorname{Pr}(>\|t\|) \\ 181.062\end{array}$ |  |
| $\begin{array}{llllll}\text { as.factor(rs174548) } 1 & 6.802 & 2.321 & 2.930 & 0.00358\end{array}$ |  |
| $\begin{array}{lllllll}\text { as.factor (rs174548)2 } & 5.438 & 4.540 & 1.198 & 0.23167\end{array}$ |  |
| --- |  |
| Residual standard error: 21.93 on 397 degrees of freedom Multiple R-squared: 0.0221, Adjusted R-squared: 0.01718 F-statistic: 4.487 on 2 and 397 DF, p-value: 0.01184 |  |
|  |  |
|  |  |
| > anova(fit1.1) |  |
| Analysis of Variance Table |  |
| Response: chol Df Sum Sq Mean Sq F value $\operatorname{Pr}(>F)$ |  |
|  |  |
| Residuals 397190875481 |  |
| --- |  |

## ANOVA: One-Way Model


Residual standard error: 21.93 on 397 degrees of freedom Multiple R-squared: 0.0221, Adjusted R-squared: 0.01718 F-statistic: 4.487 on 2 and 397 DF, p-value: 0.01184
> anova(fit1.1)
Analysis of Variance Table
Response: chol
as.factor(rs174548) Df Sum Sq Mean Sq F value Pr(>F) $\begin{array}{lll}\text { Residuals } & 397 & 190875\end{array}$

Interpretation:

- Estimated mean cholesterol for $\mathrm{C} / \mathrm{C}$ group: $181.062 \mathrm{mg} / \mathrm{dl}$
- Estimated difference in mean cholesterol levels between C/G and C/C groups: $6.802 \mathrm{mg} / \mathrm{dl}$
- Estimated difference in mean cholesterol levels between G/G and C/C groups: $5.438 \mathrm{mg} / \mathrm{dl}$



## ANOVA: One-Way Model

Alternative form:
(better if you will perform multiple comparisons)

```
> fit1.2 = lm(chol ~ -1 + as.factor(rs174548))
> summary(fit1.2)
Call:
lm(formula = chol ~ -1 + as.factor(rs174548))
Residuals:
\begin{tabular}{rrrrr} 
Residuals: & 1Q & Median & \(3 Q\) & Max \\
Min & -64.06167 & -15.91338 & -0.06167 & 14.93833 \\
-69.13605
\end{tabular}
Coefficients: Estimate Std. Error t value Pr (>|t|
as.factor(rs174548)0 181.062 1.455 124.41 <2e-16 ***
as.factor(rs174548)1 187.864 1.809 103.88 <2e-16 ***
as.factor(rs174548)2 186.500 4.300 43.37 <2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
Residual standard error: 21.93 on 397 degrees of freedom
Multiple R-squared: 0.9861, Adjusted R-squared: 0.986
F-statistic: }9383\mathrm{ on }3\mathrm{ and 397 DF, p-value: < 2.2e-16
> anova(fit1.2)
Analysis of Variance Table
Response: chol
Df Sum Sq Mean Sq F value Pr(>F)
as.factor(rs174548) 3 13534205 4511402 9383.2 < 2.2e-16 ***
Residuals 397 190875 481
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ', 1
```


## ANOVA: One-Way Model

## Alternative form:

- Different command!


## ANOVA: One-Way Model

How about this one? How is rs174548 being treated now?

Compare model fit results from (fit1.1 \& fit2).

```
> fit2 = lm(chol ~ rs174548)
> summary(fit2)
Call:
lm(formula = chol ~ rs174548)
Residuals:
\begin{tabular}{rrrrr} 
Min & \(1 Q\) & Median & \(3 Q\) & Max \\
-64.575 & -16.278 & -0.575 & 15.120 & 60.722
\end{tabular}
Coefficients:
Estimate Std. Error t value Pr (>|t|)
#rntercept) 181.575 1.411 128.723 < 2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '\star' 0.05 '.' 0.1 ', 1
Residual standard error: 21.95 on 398 degrees of freedom
Multiple R-squared: 0.01723, Adjusted R-squared: 0.01476
F-statistic: 6.977 on 1 and 398 DF, p-value: 0.008583
> anova(fit2)
Analysis of Variance Table
Response: chol
rs174548 Df Sum Sq Mean Sq F value Pr(>F)
Residuals 398 191827 482
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' }119
```



## ANOVA: One-Way Model



Residual standard error: 21.93 on 397 degrees of freedom Multiple R-squared: 0.0221 , Adjusted R-squared: 0.01718 F-statistic: 4.487 on 2 and 397 DF, p-value: 0.01184
> anova(fit1.1)
Analysis of Variance Table
Response: chol
as.factor(rs174548) Dr Sum Mean Sq F value Pr (>F) Residuals

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We rejected the null hypothesis that the mean cholesterol levels are the same across groups defined by rs174548 ( $p=0.01184$ ).

- What are the groups with differences in means?

MULTIPLE COMPARISONS

## ANOVA

MULTIPLE COMPARISONS

## ANOVA: One-Way Model

- What are the groups with differences in means?


## MULTIPLE COMPARISONS:

$$
\left.\begin{array}{r}
\mu_{0}=\mu_{1} ? \\
\mu_{0}=\mu_{2} ? \\
\mu_{1}=\mu_{2} ?
\end{array}\right\} \text { Pairwise comparisons }
$$

## Multiple Comparisons: Family-wise error rates

- Illustrating the multiple comparison problem
- Truth: null hypotheses
- Tests: pairwise comparisons - each at the 5\% level.

What is the probability of rejecting at least one?

| \#groups <br> K K | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| \#pairwise <br> comparisons <br> K K(K-1)/2 | 1 | 3 | 6 | 10 | 15 | 21 | 28 | 36 | 45 |
| P(at least <br> one sig) <br> $=1-(1-0.05)$ c | 0.05 | 0.143 | 0.265 | 0.401 | 0.537 | 0.659 | 0.762 | 0.842 | 0.901 |

That is, if you have three groups and make pairwise comparisons, each at the 5\% level, your familywise error rate (probability of making at least one false rejection) is over $14 \%$ !

Need to address this issue!
Several methods!!!

## Multiple Comparisons

- Several methods:
- None (no adjustment)
- Bonferroni
- Holm
- Hochberg
- Hommel

Available in $R$

- BH
- BY
- FDR
- ...


## Multiple Comparisons

Bonferroni adjustment: for k tests performed, use level $\mathrm{a} / \mathrm{k}$ (or multiply $P$-values by k ).

- Simple
- Conservative
- Must decide on number of tests beforehand
- Widely applicable
- Can be done without software!



## Multiple Comparisons

```
> ## -- third, adjust the p-values (or not) for multiple comparisons
> summary(mc, test=adjusted("none"))
    Simultaneous Tests for General Linear Hypotheses
Multiple Comparisons of Means: Tukey Contrasts
Fit: lm(formula = chol ~ -1 + as.factor(rs174548))
Linear Hypotheses:
Estimate Std. Error t value Pr(>|t|)
1-0== 0 6.802 2.321 2.930 0.00358 **
2 - 0 == 0 5.438 4.540 1.198 0.23167
2-1 == 0 -1.364 4.665 -0.292 0.77015
---
(Adjusted p values reported -- none method)
```


## Multiple Comparisons

```
> summary(mc, test=adjusted("bonferroni"))
    Simultaneous Tests for General Linear Hypotheses
Multiple Comparisons of Means: Tukey Contrasts
Fit: lm(formula = chol ~ -1 + as.factor(rs174548))
Linear Hypotheses:
Estimate Std. Error t value Pr(>|t|)
1-0== 0 6.802 2.321 2.930 0.0107 *
2 - 0 == 0 5.438 4.540 1.198 0.6950
2-1 == 0 -1.364 4.665 -0.292 1.0000
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ', 1
(Adjusted p values reported -- bonferroni method)
```


## Multiple Comparisons

- What if nonpairwise comparison?
- Suppose you want to compare the mean cholesterol among those with genotype $\mathrm{C} / \mathrm{C}$ with the mean cholesterol for the combined group with genotypes $\mathrm{C} / \mathrm{G}$ and $\mathrm{G} / \mathrm{G}$.

$$
\mu_{0}=\left(\mu_{1}+\mu_{2}\right) / 2
$$

Or equivalently,

$$
2 \mu_{0}=\left(\mu_{1}+\mu_{2}\right)
$$

Or equivalently,

$$
2 \mu_{0}-\mu_{1}-\mu_{2}=0
$$

## Multiple Comparisons

- What if nonpairwise comparison?
- Your turn: Suppose you want to compare the mean cholesterol among those with genotype C/G with the mean cholesterol for the combined group with genotypes $\mathrm{C} / \mathrm{C}$ and $\mathrm{G} / \mathrm{G}$.


## Multiple Comparisons

- What if nonpairwise comparison?
- Your turn: Suppose you want to compare the mean cholesterol among those with genotype $\mathrm{C} / \mathrm{G}$ with the mean cholesterol for the combined group with genotypes $\mathrm{C} / \mathrm{C}$ and $\mathrm{G} / \mathrm{G}$.

$$
\left(\mu_{0}+\mu_{2}\right) / 2=\mu_{1}
$$

Or equivalently,

$$
\mu_{0}+\mu_{2}=2 \mu_{1}
$$

Or equivalently,

$$
\mu_{0}-2 \mu_{1}+\mu_{2}=0
$$

## Multiple Comparisons

Using R for multiple comparisons with "user-defined" contrasts:

```
> contr = rbind("mean(C/G+G/G) - mean(C/C)" = c(-2, 1, 1))
> mc2 = glht(fit1, linfct =contr)
> summary(mc2, test=adjusted("none"))
    Simultaneous Tests for General Linear Hypotheses
Fit: lm(formula = chol ~ -1 + as.factor(rs174548))
Linear Hypotheses:
mean(C/G+G/G) - mean (C/C) == 0 Estimate Std. Error t value Pr(>| t|)
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
(Adjusted p values reported -- none method)
```


## Multiple Comparisons

> \#\# more than one contrast (again user-defined)
$>$ contr2 $=\operatorname{rbind}(" \operatorname{mean}(C / G+G / G)-\operatorname{mean}(C / C) "=c(-2,1,1)$,
$+\quad \operatorname{mean}(C / C+G / G)-\operatorname{mean}(C / G) "=c(1,-2,1))$
$>\mathrm{mc} 3=$ glht(fit1, linfct $=$ contr2)
> summary (mc3, test=adjusted("none"))
Simultaneous Tests for General Linear Hypotheses
Fit: $\operatorname{lm}(f o r m u l a=$ chol $\sim-1+$ as.factor (rs174548))
Linear Hypotheses:
Estimate Std. Error $t$ value $\operatorname{Pr}(>|t|)$

$\operatorname{mean}(\mathrm{C} / \mathrm{G}+\mathrm{G} / \mathrm{G})-\operatorname{mean}(\mathrm{C} / \mathrm{C})=$|  | 0 | 12.241 | 5.499 | 2.226 |
| ---: | ---: | ---: | ---: | ---: |


$\operatorname{mean}(C / C+G / G)-\operatorname{mean}(C / G)==$| 0 | -8.166 | 5.805 | -1.407 | 0.1603 |
| :--- | :--- | :--- | :--- | :--- | :--- |

Signif. codes: $0{ }^{\prime * * * '} 0.001$ '**' 0.01 '*' 0.05 '.' 0.1 ', 1
(Adjusted p values reported -- none method)
> summary(mc3, test=adjusted("bonferroni"))
Simultaneous Tests for General Linear Hypotheses
Fit: lm(formula = chol $\sim-1+$ as.factor(rs174548))
Linear Hypotheses:
Estimate Std. Error $t$ value $\operatorname{Pr}(>|t|)$

$\operatorname{mean}(\mathrm{C} / \mathrm{G}+\mathrm{G} / \mathrm{G})-\operatorname{mean}(\mathrm{C} / \mathrm{C})==$| 0 | 12.241 | 5.499 | 2.226 | 0.0531 |
| :--- | :--- | :--- | :--- | :--- | :--- |

$\operatorname{mean}(\mathrm{C} / \mathrm{C}+\mathrm{G} / \mathrm{G})-\operatorname{mean}(\mathrm{C} / \mathrm{G})==\begin{array}{llllll}0 & -8.166 & 5.805 & -1.407 & 0.3205\end{array}$
---
Signif. codes: $0{ }^{\prime * * * ’} 0.001$ '**' 0.01 ‘*' 0.05 '. 0.1 ', 1
(Adjusted p values reported -- bonferroni method)

## Multiple Comparisons

- What about using other adjustment methods?
- For example, we used:
> summary(mc, test=adjusted("bonferroni"))
(all pairwise comparisons, with Bonferroni adjustment)
- Other options, in place of "bonferroni", are:
- summary(mc, test=adjusted("holm"))
- summary(mc, test=adjusted("hochberg"))
- summary(mc, test=adjusted("hommel"))
- summary(mc, test=adjusted("BH"))
- summary(mc, test=adjusted("BY"))
- summary(mc, test=adjusted("fdr"))

Results, in this particular example, are basically the same, but they don't need to be! Different criteria could lead to different results!

## Multiple Comparisons

```
> summary(mc, test=adjusted("fdr"))
    Simultaneous Tests for General Linear Hypotheses
Multiple Comparisons of Means: Tukey Contrasts
Fit: lm(formula = chol ~ -1 + as.factor(rs174548))
Linear Hypotheses:
Estimate Std. Error t value Pr(>|t|)
1-0 == 0 6.802 2.321 2.930 0.0107 *
2-0 == 0 5.438 4.540 1.198 0.3475
2-1 == 0 -1.364 4.665 -0.292 0.7702
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
(Adjusted p values reported -- fdr method)
```


## Multiple Comparisons

- FDR (False Discovery Rate)
- Less conservative procedure for multiple comparisons
- Among rejected hypotheses, FDR controls the expected proportion of incorrectly rejected null hypotheses (that is, type I errors).



## ANOVA Assumptions

- Recall the assumptions for classical ANOVA are:


## Independence

Normality
Equal variance


## Bartlett's test

- We assume that variances are the same across populations
- Bartlett's test allows you to test the hypothesis that the population variances are the same (versus they are not all equal).

```
> bartlett.test(chol ~ as.factor(rs174548))
    Bartlett test of homogeneity of variances
data: chol by as.factor(rs174548)
Bartlett's K-squared = 4.8291, df = 2, p-value = 0.0894
```


## Bartlett' s test?

- No real need to test variances!
- You can perform one-way ANOVA allowing for unequal variances!
- You can perform one-way ANOVA - using the regression framework with robust standard errors!


## One-Way ANOVA with robust standard errors

```
> summary(gee(chol ~ as.factor(rs174548), id=seq(1,length(chol))))
    Beginning Cgee S-function, @(#) geeformula.q 4.13 98/01/27
    running glm to get initial regression estimate
    (Intercept) as.factor(rs174548)1 as.factor(rs174548)2
    181.061674 6.802272 5.438326
    GEE: GENERALIZED LINEAR MODELS FOR DEPENDENT DATA
    gee S-function, version 4.13 modified 98/01/27 (1998)
Model:
    Link: Identity
    Variance to Mean Relation: Gaussian
    Correlation Structure: Independent
    Cal1
    gee(formula = chol ~ as.factor(rs174548), id = seq(1, length(chol)))
Summary of Residuals:
    Min 1Q
-64.06167401 -15.91337769 -0.06167401 14.93832599
Coefficients:
                                    Naive S.E. Naive z Robust S.E. Robust z
(Intercept) 181.061674 1.455346 124.411431 1.400016 129.328297
as.factor(rs174548)1 
as.factor(rs174548)2 
Estimated Scale Parameter: 480.7932
Number of Iterations: 1
```


## Kruskal-Wallis Test

- Non-parametric analogue to the one-way ANOVA
- Based on ranks


## - In our example:

```
> kruskal.test(chol ~ as.factor(rs174548))
    Kruskal-Wallis rank sum test
data: chol by as.factor(rs174548)
Kruskal-Wallis chi-squared = 7.4719, df = 2, p-value = 0.02385
```

- Conclusion:
- Evidence that the cholesterol distribution is not the same across all groups.
- With the global null rejected, you can also perform pairwise comparisons [Wilcoxon rank sum], but adjust for multiplicities!


## Multiple Comparisons

```
(following Kruskal-Wallis Test)
> wilcox.test(chol[rs174548!=0] ~rs174548[rs174548!=0])
Wilcoxon rank sum test with continuity correction
data: chol[rs174548 ! = 0] by rs174548[rs174548 != 0]
\(\mathrm{W}=1974.5\), p -value \(=0.789\)
alternative hypothesis: true location shift is not equal to 0
> wilcox.test(chol[rs174548!=1] ~rs174548[rs174548!=1])
Wilcoxon rank sum test with continuity correction
data: chol[rs174548 != 1] by rs174548[rs174548 != 1]
\(W=2482, \mathrm{p}\)-value \(=0.1849\)
alternative hypothesis: true location shift is not equal to 0
> wilcox.test(chol[rs174548!=2] ~rs174548[rs174548!=2])
Wilcoxon rank sum test with continuity correction
data: chol[rs174548 != 2] by rs174548[rs174548 != 2]
\(\mathrm{W}=14025.5\), p -value \(=0.009221\)
alternative hypothesis: true location shift is not equal to 0
```



Two-way ANOVA models

ANOVA: Two-Way Model
Motivation:

- Scientific question:
- Assess the effect of rs174548 and gender on cholesterol levels.


## ANOVA: Two-Way Model

- Factors: A and B
- Goals:
- Test for main effect of $A$
- Test for main effect of B
- Test for interaction effect of $A$ and $B$


## ANOVA: Two-Way Model

- To simplify discussion, assume that factor A has three levels, while factor B has two levels




## ANOVA: Two-Way Model

- Recall:
- Categorical variables can be represented with "dummy" variables
- Interactions are represented with "cross-products"


## ANOVA: Two-Way Model

- Model 1:
$E\left[Y \mid A_{2}, A_{3}, B_{2}\right]=\beta_{0}+\beta_{1} A_{2}+\beta_{2} A_{3}+\beta_{3} B_{2}$.
- What are the means in each combination-group?

|  | $A_{1}$ | $A_{2}$ | $A_{3}$ |
| :--- | :--- | :--- | :--- |
| $B_{1}$ | $\mu_{11}=\beta_{0}$ | $\mu_{21}=\beta_{0}+\beta_{1}$ | $\mu_{31}=\beta_{0}+\beta_{2}$ |
| $B_{2}$ | $\mu_{12}=\beta_{0}+\beta_{3}$ | $\mu_{22}=\beta_{0}+\beta_{1}+\beta_{3}$ | $\mu_{32}=\beta_{0}+\beta_{2}+\beta_{3}$ |

## ANOVA: Two-Way Model

- Model 1:
$E\left[Y \mid A_{2}, A_{3}, B_{2}\right]=\beta_{0}+\beta_{1} A_{2}+\beta_{2} A_{3}+\beta_{3} B_{2}$.

|  | $A_{1}$ | $A_{2}$ | $A_{3}$ |
| :--- | :--- | :--- | :--- |
| $B_{1}$ | $\mu_{11}=\beta_{0}$ | $\mu_{21}=\beta_{0}+\beta_{1}$ | $\mu_{31}=\beta_{0}+\beta_{2}$ |
| $B_{2}$ | $\mu_{12}=\beta_{0}+\beta_{3}$ | $\mu_{22}=\beta_{0}+\beta_{1}+\beta_{3}$ | $\mu_{32}=\beta_{0}+\beta_{2}+\beta_{3}$ |

Model with no interaction:
-Difference in means between groups defined by factor $B$ does not depend on the level of factor A .
-Difference in means between groups defined by factor A does not depend on the level of factor B.

## ANOVA: Two-Way Model

- Model 2:
$E\left[Y \mid A_{2}, A_{3}, B_{2}\right]=\beta_{0}+\beta_{1} A_{2}+\beta_{2} A_{3}+\beta_{3} B_{2}+\beta_{4} A_{2} B_{2}+\beta_{5} A_{3} B_{2}$
- What are the means in each combination-group?

|  | $A_{1}$ | $A_{2}$ | $A_{3}$ |
| :--- | :--- | :--- | :--- |
| $B_{1}$ | $\mu_{11}=\beta_{0}$ | $\mu_{21}=\beta_{0}+\beta_{1}$ | $\mu_{31}=\beta_{0}+\beta_{2}$ |
| $B_{2}$ | $\mu_{12}=\beta_{0}+\beta_{3}$ | $\mu_{22}=\beta_{0}+\beta_{1}+\beta_{3}+\beta_{4}$ | $\mu_{32}=\beta_{0}+\beta_{2}+\beta_{3}+\beta_{5}$ |

## ANOVA: Two-Way Model

- Three (possible) tests
- Interaction of $A$ and $B$ (may want to start here)
- Rejection would imply that differences between means of A depends on the level of B (and vice-versa) so stop
- Main effect of $A$
- Test only if no interaction
- Main effect of B
- Test only if no interaction
[ Note: If you have one observation per cell, you cannot test interaction! ]


## ANOVA: Two-Way Model

- Model without interaction
$E\left[Y \mid A_{2}, A_{3}, B_{2}\right]=\beta_{0}+\beta_{1} A_{2}+\beta_{2} A_{3}+\beta_{3} B_{2}$.

How do we test for main effect of factor $A$ ?
$\mathrm{H}_{0}: \beta_{1}=\beta_{2}=0$ vs. $\mathrm{H}_{1}: \beta_{1}$ or $\beta_{2}$ not zero

How do we test for main effect of factor $B$ ?
$H_{0}: \beta_{3}=0$ vs. $H_{1}: \beta_{3}$ not zero

## ANOVA: Two-Way Model

- Model with interaction:

$$
E\left[Y \mid A_{2}, A_{3}, B_{2}\right]=\beta_{0}+\beta_{1} A_{2}+\beta_{2} A_{3}+\beta_{3} B_{2}+\beta_{4} A_{2} B_{2}+\beta_{5} A_{3} B_{2}
$$

How do we test for interactions?
$H_{0}: \beta_{4}=\beta_{5}=0$ vs.
$H_{1}: \beta_{4}$ or $\beta_{5}$ not zero

IMPORTANT:
If you reject the null, do not test main effects!!!

## ANOVA: Two-Way Model (without interaction)

```
> fit1 = lm(chol ~ as.factor(sex) + as.factor(rs174548))
> summary(fit1)
    lm(formula = chol ~ as.factor(sex) + as.factor(rs174548))
    Residuals:
    Mrrrrr
    Coefficients:
    # Estimate Std. Error t value Pr (>|t|)
    as.factor(sex)1 11.053 1.126 5.199 3.22e-07 ***
    as.factor(rs174548)1 7.236 2.250 3.215 0.00141 *
    as.factor(rs174548)2 5.184 4.398 1.179 0.23928
    Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ', 1
    Residual standard error: 21.24 on 396 degrees of freedom
    Multiple R-squared: 0.08458, Adjusted R-squared: 0.07764
    F-statistic: 12.2 on 3 and 396 DF, p-value: 1.196e-07
    > anova(fit0,fit1)
    Analysis of Variance Table
    Model 1: chol ~ as.factor(sex)
    Model 2: chol ~ as.factor(sex) + as.factor(rs174548)
    Res.Df RSS Df Sum of Sq F Pr(>F)
1 398 183480
    2 396 178681 2 4799.1 5.318 0.005259 **
```


## ANOVA: Two-Way Model (without interaction)

- Interpretation of results:


## $>$ fit1 $=1 \mathrm{~m}($ chol $\sim$ as.factor (sex) + as.factor (rs174548))

 $>$ summary (fit1)Call:
lm(formula $=$ chol $\sim$ as.factor(sex) + as.factor(rs174548))
Residuals:
$\begin{array}{rrrrr}\text { Min } & 10 & \text { Median } & 39 & \text { Max } \\ -66.6534 & -14.4633 & -0.6008 & 15.4450 & 57.6350\end{array}$
Coefficients:

|  | Estimate | Std. Error | t value | $\operatorname{Pr}(>\|t\|)$ |  |
| :--- | ---: | ---: | ---: | ---: | ---: | :--- |
| (Intercept) | 175.365 | 1.786 | 98.208 | $<2 \mathrm{e}-16$ | *** |
| as.factor(sex)1 | 11.053 | 2.126 | 5.199 | $3.22 \mathrm{e}-07$ | *** |
| as.factor(rs174548)1 | 7.236 | 2.250 | 3.215 | 0.00141 | ** |
| as.factor(rs174548)2 | 5.184 | 4.398 | 1.179 | 0.23928 |  |

$\begin{array}{llllll}\text { as.factor(rs174548)2 } & 5.184 & 4.398 & 1.179 & 0.23928\end{array}$
Residual standard error: 21.24 on 396 degrees of freedom Multiple R-squared: 0.08458 , Adjusted R-squared: 0.07764 F-statistic: 12.2 on 3 and 396 DF , p-value: $1.196 \mathrm{e}-07$
> anova(fit0,fit1)
Analysis of Variance Table
Model 1: chol ~ as.factor(sex)
Model 2: chol ~ as.factor(sex) + as.factor(rs174548)
Res.Df RSS Df Sum of $\mathrm{Sq} \quad \mathrm{F} \quad$ Pr $(>\mathrm{F})$
1398183480
$2396178681 \quad 2 \quad 4799.1 \quad 5.318 \quad 0.005259$ **

- Estimated mean cholesterol for male C/C group: $175.37 \mathrm{mg} / \mathrm{dl}$
- Estimated difference in mean cholesterol levels between females and males adjusted by genotype: $11.053 \mathrm{mg} / \mathrm{dl}$
- Estimated difference in mean cholesterol levels between C/G and C/C groups adjusted by gender: $7.236 \mathrm{mg} / \mathrm{dl}$
- Estimated difference in mean cholesterol levels between $\mathrm{G} / \mathrm{G}$ and C/C groups adjusted by gender: $5.184 \mathrm{mg} / \mathrm{dl}$
- There is evidence that cholesterol is associated with gender ( $p<0.001$ ).
- There is evidence that cholesterol is associated with genotype ( $p=0.005$ )


## ANOVA: Two-Way Model (without interaction)

- In words:
- Adjusting for sex, the difference in mean cholesterol comparing C/G to C/C is 7.236 and comparing $\mathrm{G} / \mathrm{G}$ to $\mathrm{C} /$ C is 5.184 .
- This difference does not depend on sex
- (this is because the model does not have an interaction between sex and genotype!)


## ANOVA: Two-Way Model (with interaction)

```
> fit2 = lm(chol ~ as.factor(sex) * as.factor(rs174548))
> summary(fit2)
Call:
lm(formula = chol ~ as.factor(sex) * as.factor(rs174548))
Residuals:
    Min 1Q Median 3Q Max
-70.5286 -13.6037 -0.9736 14.1709 54.8818
Coefficients:
(Intercept) 178.1182 2.0089 88.666 < 2e-16 ***
as.factor(sex)1 5.7109 2.7982 2.041 0.04192 *
as.factor(rs174548)1 0.9597 3.1306 0.307 0.75933
as.factor(rs174548)2 -0.2015 6.4053 -0.031 0.97492
as.factor(sex)1:as.factor(rs174548)1 12.7398 4.4650 2.853 0.00456 **
as.factor(sex) 1:as.factor(rs174548)2 10.2296 
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
Residual standard error: 21.07 on 394 degrees of freedom
Multiple R-squared: 0.1039, Adjusted R-squared: 0.09257
F-statistic: 9.14 on 5 and 394 DF, p-value: 3.062e-08
```


## ANOVA: Model comparison

```
> anova(fit1,fit2)
Analysis of Variance Table
Model 1: chol ~ as.factor(sex) + as.factor(rs174548)
Model 2: chol ~ as.factor(sex) * as.factor(rs174548)
    Res.Df RSS Df Sum of Sq F Pr(>F)
1 396 178681
2 394 174902 2 3779 4.2564 0.01483 *
---
Signif. codes: 0 '***' 0.001 '**' 0.01 '*’ 0.05 '.' 0.1 ', 1
```




## ANCOVA MODELS

(aka ANACOVA)

ANalysis of COVAriance Models (ANCOVA) Motivation:

- Scientific question:
- Assess the effect of rs174548 on cholesterol levels adjusting for age

ANalysis of COVAriance Models (ANCOVA)

- ANOVA with one or more continuous variables
- Equivalent to regression with "dummy" variables and continuous variables
- Primary comparison of interest is across $k$ groups defined by a categorical variable, but the $k$ groups may differ on some other potential predictor or confounder variables [also called covariates].


## ANalysis of COVAriance Models (ANCOVA)

- To facilitate discussion assume
- Y: continuous response (e.g. cholesterol)
- X: continuous variable (e.g. age)
- Z: dummy variable (e.g. indicator of C/G or G/G versus C/C)
- Model: $Y=\beta_{0}+\beta_{1} X+\beta_{2} Z+\beta_{2} X Z+\varepsilon$

Note that:

$$
\begin{aligned}
& Z=0 \Rightarrow E[Y \mid X, Z=0]=\beta_{0}+\beta_{1} X \\
& Z=1 \Rightarrow E[Y \mid X, Z=1]=\left(\beta_{0}+\beta_{2}\right)+\left(\beta_{1}+\beta_{3}\right) X
\end{aligned}
$$

This model allows for different intercepts/slopes for each group.

## ANCOVA

- Testing coincident lines: $H_{0}: \beta_{2}=0, \beta_{3}=0$
- Compares overall model with reduced model

$$
Y=\beta_{0}+\beta_{1} X+\varepsilon
$$

- Testing parallelism: $H_{0}: \beta_{3}=0$
- Compares overall model with reduced model

$$
Y=\beta_{0}+\beta_{1} X+\beta_{2} Z+\varepsilon
$$

## ANCOVA

```
> fit0 = lm(chol ~ as.factor(rs174548))
> summary(fit0)
Call:
lm(formula = chol ~ as.factor(rs174548))
Residuals: 
Coefficients:
lrrercept) 
lllll
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 ', 0.1 ', 1
Residual standard error: 21.93 on 397 degrees of freedom
Multiple R-squared: 0.0221, Adjusted R-squared: 0.01718
F-statistic: 4.487 on 2 and 397 DF, p-value: 0.01184
> anova(fit0)
Analysis of Variance Table
Response: chol
S Df Sum Sq Mean Sq F value Pr(>F)
```



```
---
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ', 1
```






## ANCOVA

- In summary:
- If the slopes are not equal, then age is an effect modifier

$$
\begin{gathered}
E[Y \mid x, z]=\beta_{0}+\beta_{1} x+\beta_{2}(C G)+\beta_{3}(G G)+ \\
\beta_{4}(x * C G)+\beta_{5}(x * G G)
\end{gathered}
$$

- If the slopes are the same,

$$
E[Y \mid x, z]=\beta_{0}+\beta_{1} x+\beta_{2}(C G)+\beta_{3}(G G)
$$

## ANCOVA

- If the slopes are the same,

$$
E[Y \mid x, z]=\beta_{0}+\beta_{1} x+\beta_{2}(C G)+\beta_{3}(G G)
$$

- then one can obtain adjusted means for the three genotypes using the mean age over all groups
- For example, the adjusted means for the three groups would be

$$
\begin{aligned}
& \overline{\mathrm{Y}}_{1}(\mathrm{adj})=\hat{\beta}_{0}+\bar{x} \hat{\beta}_{1} \\
& \overline{\mathrm{Y}}_{2}(\mathrm{adj})=\left(\hat{\beta}_{0}+\hat{\beta}_{2}\right)+\bar{x} \hat{\beta}_{1} \\
& \overline{\mathrm{Y}}_{3}(\mathrm{adj})=\left(\hat{\beta}_{0}+\hat{\beta}_{3}\right)+\bar{x} \hat{\beta}_{1}
\end{aligned}
$$

## ANCOVA

```
## unadjusted mean cholesterol levels for different genotypes
> predict(fit0, new=data.frame(rs174548=0))
181.0617
> predict(fit0, new=data.frame(rs174548=1))
187.8639
> predict(fit0, new=data.frame(rs174548=2))
1
186.5
> ## mean cholesterol for different genotypes adjusted by age
> predict(fit1, new=data.frame(age=mean(age),rs174548=0))
1
180.9013
> predict(fit1, new=data.frame(age=mean(age),rs174548=1))
88.2026
> predict(fit1, new=data.frame(age=mean(age),rs174548=2))
1
185.9856
```



## Experimental Designs \& ANOVA

- This section is not intended to be comprehensive
- No endorsement for any of the articles cited here


## Tool Kit

## - Controls and Placebos:

- Provides a baseline comparison with test groups


## - Blinding:

- When successfully applied, it eliminates the possibility that the end comparison measures expectations rather than real treatment differences
- Blocking:
- Arranges units into homogeneous subgroups so that treatments can be randomly assigned to units within each block
- Improves precision for treatment comparisons
- Controls for confounding variables by grouping experimental units into blocks with similar values of the variable


## Tool Kit

## - Stratification

- Involves partitioning of population units into homogeneous subgroups - called strata - and performing random sampling of population units in each strata
- (stratification pertains to random sampling; blocking pertains to random assignment)
- Covariates
- Inclusion may control for potentially confounding factors
- Inclusion may improve precision in treatment comparisons
- Randomization
- Allows for controlling for factors not explictly controlled for in the design (by blocking) or in the analysis (by covariates)
- Enables causal inferences


## Tool Kit

- Random Sampling
- Means employing a random procedure to select units from a population
- To ensure that sample is representative of the population
- To permit an inference that patterns observed in the sample are characteristic of patterns in the population as a whole
- Replication
- It refers to assigning one treatment to multiple units within each block.
- Increases precision for treatment effects (increased sample size)
- Allows for model assessment
- Balance
- Same number of units to each treatment
- Optimizes precision for treatment comparisons


## Terminology

- Treatments
- A factor level in a single-factor study or a combination of factor levels in a multi-factor study
- How many factors should be examined?
- How many levels should each factor have?
- Experimental units
- Smallest unit of the experiment such that any two different experimental units may receive different treatments


## One-Way Data Patterns

Factor


Factor: categorical predictor
Two-Way Data Patterns

| Y | Y | Y |
| :---: | :---: | :---: |
| Y | Y | Y |
| $Y$ | Y | Y |
| $Y$ | Y | Y |

Single observation per cell

| Factor 1 |  |  |
| :---: | :---: | :---: |
| YYY | YYY | YYY |
| YYY | YYY | YYY |
| \% YYY | YYY | YYY |
| YYY | YYY | YYY |

Equal replication per cell

| YY | YYY | YYYYYY |
| :---: | :---: | :---: |
| YYY | YYYY | YY |
| Y | YYY | YYYY |
| YYYYY | YY | Y |

Non-systematic replications

## Completely Randomized Design

- Treatments are allocated to the experimental units completely at random
- Every experimental unit has an equal chance of receiving any of the treatments
- Simple \& flexible
- Allows for any number of treatments
- Sample sizes can vary from treatment to treatment
- Inefficient when the experimental units are heterogeneous


## Completely Randomized Design



## Completely Randomized Design: an Example

- Title: "Hepatocyte growth factor incorporated chitosan nanoparticles augment the differentiation of stem cell into hepatocytes for the recovery of liver cirrhosis in mice."
- Authors: Pulavendran S, Rose C, Mandal AB. J Nanobiotechnology. 2011 Apr 28;9:15.
- Abstract [partial]:
- background: Short half-life and low levels of growth factors in the niche of injured microenvironment necessitates the exogenous and sustainable delivery of growth factors along with stem cells to augment the regeneration of injured tissues.
- METHODS: Recombinant human hepatocyte growth factor (HGF) was incorporated into chitosan nanoparticles (CNP) by ionic gelation method and studied for its morphological and physiological characteristics. Cirrhotic mice received either hematopoietic stem cells (HSC) or mesenchymal stemcells (MSC) with or without HGF incorporated chitosan nanoparticles (HGF-CNP) and saline as control. Biochemical, histological, immunostaining and gene expression assays were carried out using serum and liver tissue samples [...].
- results: Serum levels of selected liver protein and enzymes were significantly increased in the combination of MSC and HGF-CNP (MSC+HGF-CNP) treated group.
- conclusion: [...] Transplantation of bone marrow MSC in combination with HGF-CNP could be an ideal approach for the treatment of liver cirrhosis.


## Completely Randomized Design: Exercise

What is the goal of the experiment?

What is(are) the response variables?

- What are the factors?
- How many levels?
- Statistical model?


## Factorial Design

- A factorial design is used to evaluate two or more factors simultaneously.
- Factorial designs are more efficient than one-factor-at-a-time designs
- Factorial designs allow for investigations of interactions.


## Factorial Design: an example

- Title: "Fermentable fiber ameliorates fermentable protein-induced changes in microbial ecology, but not the mucosal response, in the colon of piglets".
- Pieper R, Kröger S, Richter JF, Wang J, Martin L, Bindelle J, Htoo JK, von Smolinski D, Vahjen W, Zentek J, Van Kessel AG. J Nutr. 2012 Apr;142(4):661-7. Epub 2012 Feb 22.
- Abstract (partial): Dietary inclusion of fermentable carbohydrates (fCHO) is reported to reduce large intestinal formation of putatively toxic metabolites derived from fermentable proteins (fCP). However, the influence of diets high in fCP concentration on epithelial response and interaction with fCHO is still unclear. Thirty-two weaned piglets were fed 4 diets in a $\mathbf{2 \times 2} \mathbf{2}$ factorial design with low fCP/low fCHO [14.5\% crude protein (CP)/ $14.5 \%$ total dietary fiber (TDF)]; low fCP/high fCHO (14.8\% CP/16.6\% TDF); high fCP low fCHO ( $19.8 \%$ CP/14.5\% TDF); and high fCP/high fCHO ( $20.1 \%$ CP/18.0\% TDF) as dietary treatments. After 21-23 d, pigs were killed and colon digesta and tissue samples analyzed for indices of microbial ecology, tissue expression of genes for cell turnover, cytokines, mucus genes (MUC), and oxidative stress indices. Pig performance was unaffected by diet. [...] High dietary fCP increased ( $\mathrm{P}<0.05$ ) expression of PCNA, IL1 $\beta$, IL10, TGFß, MUC1, MUC2, and MUC20, irrespective of fCHO concentration.


## Factorial Design: Exercise <br> - What is the goal of the experiment?

- What is(are) the response variables?
- What are the factors?
- For each factor, how many levels?
- How many treatments?
- Statistical model?


## Factorial Design: an example

TABLE 3 Relative mRNA abundance of proliferating cell nuclear antigen, caspase 3, pro- and antiinflammatory cytokines, and mucus genes in the colon of piglets fed diets containing a low or high concentration of fCHO or $\mathrm{fCP}^{1,2}$

| Gene | Low fCP |  | High fCP |  | $P$ value ${ }^{3}$ |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Low fCHO | High fCHO | Low fCH0 | High fCHO | fCHO | fCP | fCHO x fCP |
| PCNA | $0.81 \pm 0.05$ | $0.79 \pm 0.04$ | $0.89 \pm 0.08$ | $0.90 \pm 0.04$ | 0.94 | $<0.05$ | 0.76 |
| CASP | $0.80 \pm 0.04$ | $0.85 \pm 0.06$ | $0.88 \pm 0.06$ | $0.85 \pm 0.04$ | 0.83 | 0.46 | 0.37 |
| LL1 $\beta$ | $0.87 \pm 0.11$ | $0.89 \pm 0.07$ | $1.01 \pm 0.10$ | $1.05 \pm 0.07$ | 0.71 | $<0.05$ | 0.89 |
| IL6 | $0.76 \pm 0.13$ | $0.81 \pm 0.15$ | $1.04 \pm 0.19$ | $1.01 \pm 0.15$ | 0.96 | 0.07 | 0.77 |
| 1210 | $0.92 \pm 0.07$ | $0.90 \pm 0.09$ | $1.09 \pm 0.08$ | $1.05 \pm 0.04$ | 0.61 | $<0.05$ | 0.86 |
| TGFß | $0.88 \pm 0.09$ | $0.85 \pm 0.10$ | $1.11 \pm 0.09$ | $1.07 \pm 0.05$ | 0.61 | $<0.01$ | 0.93 |
| MUC1 | $0.71 \pm 0.11$ | $0.73 \pm 0.09$ | $0.89 \pm 0.09$ | $0.87 \pm 0.08$ | 0.83 | 0.05 | 0.61 |
| MUC2 | $0.84 \pm 0.14$ | $0.82 \pm 0.09$ | $1.05 \pm 0.10$ | $1.00 \pm 0.08$ | 0.97 | 0.05 | 0.79 |
| MUC20 | $0.81 \pm 0.05$ | $0.79 \pm 0.04$ | $0.89 \pm 0.08$ | $0.90 \pm 0.04$ | 0.72 | $<0.05$ | 0.85 |

${ }^{1}$ Data are mean $\pm \mathrm{SE}, n=8 /$ group. fCHO , fermentable carbohydrate; fCP , fermentable crude protein.
${ }^{2}$ Values are given as arbitrary values based on standard curves using pooled RNA samples. The mRNA abundance was normalized using $18 S$ rRNA, 60 S ribosomal protein L19 (RPL19), hypoxanthine phosphoribosyltransferase I (HPRT), and $\beta$-Actin as housekeeping genes.
${ }^{3}$ The $P$ values indicate main effects for fCP and fCHO , respectively.
Are these results unexpected?
Any concerns?

## Randomized Complete Block Designs

- Experimental units are assigned to homogeneous groups (aka "blocks").
- Reduces the variation and increases the precision of treatment comparisons
- Members of each block are randomly assigned to different treatments.
- Randomized complete block design: each block contains all treatment combinations
- Randomized incomplete block design: number of treatments exceeds the number of units in each block


## Randomized Complete Block Designs



Large N-S variability
Small E-W variability
Block 2
Within each block, a separate randomization allocates treatments to experimental units

Block 1


Block 3 $\square$ B C $\square$

## Randomized Complete Block Designs

- Factors:
- Block (control factor)
- Treatment (factor of interest)
- Statistical Model
- Two-way ANOVA model
- (additive model with single replication)


## Randomized Complete Block <br> Designs: An example

A researcher studied the effects of three experimental diets with varying fat contents on the total lipid (fat) level in plasma. Total lipid level is a widely used predictor of coronary heart disease. Fifteen male subjects who were within $20 \%$ of their ideal body weight were grouped into five blocks according to age. Within each block, the three experimental diets were randomly assigned to three subjects. Data on reduction in lipid level (in grams per liter) after the subjects were on the diet for a fixed period of time were recorded.

|  | Randomized Complete Block <br> Designs: An example |  |  |
| :--- | :--- | :--- | :--- |
|  | Fat Content of Diet |  |  |
| Age Group | Extremely Low | Fairly Low | Moderately Low |
| Ages 15-24 | 0.73 | 0.67 | 0.15 |
| Ages 25-34 | 0.86 | 0.75 | 0.21 |
| Ages 35-44 | 0.94 | 0.81 | 0.26 |
| Ages 45-54 | 1.4 | 1.32 | 0.75 |
| Ages 55-64 | 1.62 | 1.41 | 0.78 |

## Randomized Complete Block Designs: Exercise

- What is the goal of the experiment?
- What is (are) the response variables?
- What is the factor of interest? What is the blocking factor? For each factor, how many levels?
- How many treatments?
- Statistical model?


## Randomized Complete Block Designs: Another example

TITLE: UUV REPAIR AND RESISTANCE TO SOLAR UV-B IN AMPHIBIAN EGGS - A LINK TO POPULATION DECLINES"
-Author(s): BLAUSTEIN, AR (BLAUSTEIN, AR); HOFFMAN, PD (HOFFMAN, PD); HOKIT, DG (HOKIT, DG); KIESECKER, JM (KIESECKER, JM); WALLS, SC (WALLS, SC); HAYS, JB (HAYS, JB) Source: PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE UNITED STATES OF AMERICA Volume: 91 Issue: 5 Pages: 1791-1795
-Abstract [partial]: The populations of many amphibian species, in widely scattered habitats, appear to be in severe decline; other amphibians show no such declines. There is no known single cause for the declines, but their widespread distribution suggests involvement of global agents-increased UV-B radiation, for example. We addressed the hypothesis that differential sensitivity among species to UV radiation contributes to these population declines. We focused on species-specific differences in the abilities of eggs to repair UV radiation damage to DNA and differential hatching success of embryos exposed to solar radiation at natural oviposition sites. Quantitative comparisons of activities of a key UV-damage-specific repair enzyme, photolyase, among oocytes and eggs from 10 amphibian species were reproducibly characteristic for a given species but varied $>80$-fold among the species. Levels of photolyase generally correlated with expected exposure of eggs to sunlight. Among the frog and toad species studied, the highest activity was shown by the Pacific treefrog (Hyla regilla), whose populations are not known to be in decline. The Western toad (Bufo boreas) and the Cascades frog (Rana cascadae), whose populations have declined markedly, showed significantly lower photolyase levels. [...] These observations are thus consistent with the UV-sensitivity hypothesis.

## Randomized Complete Block Designs: Another example

- Goal: Is the failure rate different for species with different levels of activity of photolyase?
- Factors:
- UV-B Filter:
- UV-B blocking filter
- UV-B transmitting filter
- No Filter
- Species:
- Toad (Bufo boreas)
- Tree frog (Hyla regilla)
- Cascade frog (Rana cascadae)
- Randomization:
- Filtering treatments and egg species randomly assigned to enclosures constructed to contain clusters of 150 eggs 5


## Randomized Complete Block Designs: Another example

- Four sites: [three with single species]
- Sparks Lake (tree frog)
- Small Lake (Cascade frog)
- Lost Lake (toad)
- Three Creeks (all three species)
- Only eggs of naturally occurring species were assigned to enclosures at each site
- Blocking factor: Amphibian species/sites
- At Three Creeks: experiment is a 3 by 3 factorial design
- At other sites: single factor experiment


## Randomized Complete Block Designs



Large N-S variability
Small E-W variability

Block 3 |  | A | B | D |
| :--- | :--- | :--- | :--- |
|  |  | C |  |

## Latin Square Designs

- Employs two blocking variables ("row" and "column")
- Allows for better control of experimental variation
- Features:
- There are r treatments
- There are two blocking variables; each with r categories
- Each row and each column in the design contains all treatments
- Only one treatment per combination block


## Latin Square Designs

Latin square for 3 treatments

| $A$ | $B$ | $C$ |
| :---: | :---: | :---: |
| $C$ | $A$ | $B$ |
| $B$ | $C$ | $A$ |

Each treatment appears exactly once in each column and in each row.

Latin square for 4 treatments

| $A$ | $B$ | $D$ | $C$ |
| :---: | :---: | :---: | :---: |
| $D$ | $C$ | $A$ | $B$ |
| $B$ | $D$ | $C$ | $A$ |
| $C$ | $A$ | $B$ | $D$ |

## Latin Square Designs: An example

- In a study of chemotherapy treatments for breast cancer, researchers wanted to control for the effects of age and BMI.

|  |  | Age (years) |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | $[40,50)$ | $[50,60)$ | $[60,70)$ | 70+ |
| BMI | <20 | A | B | C | D |
|  | $[20,25)$ | B | C | D | A |
|  | $[25,30)$ | C | D | A | B |
|  | 30+ | D | A | B | C |

## Latin Square Designs: randomization

- Randomization is a bit complex because there are multiple possible Latin squares.
- Example:
- For $r=4$, there are 576 possible Latin squares (4 are of standard form).
- A Latin square is said to be in standard form (also, normalized or reduced) if both its first row and its first column are in their natural order. For example, for $\mathrm{r}=4$,

| A | B | C | D |
| :---: | :---: | :---: | :---: |
| B | C | D | A |
| C | D | A | B |
| D | A | B | C |

## Latin Square Designs: randomization

- One chooses one Latin square randomly in a particular experiment.
- This may be done by writing down any legitimate Latin square and then randomly permuting rows and columns.
- "Algorithm":
- Choose a standard Latin square (may or not be at random).
- Randomly permute all rows.
- Randomly permute all columns.
- Randomly assign treatments to the letters A, B, C, etc.

| A | B | C | D |
| :--- | :--- | :--- | :--- |
| B | C | D | A |
| R | Rows: |  |  |
| C | D | A | B |
|  | $(2,4,1,3)$ |  |  |
| D | A | B | C |


| B | C | D | A |
| :---: | :---: | :---: | :---: |
| D | A | B | C |
| A | B | C | D |
| $(3,4,4,1)$ |  |  |  |
| C | D | A | B |


| $\mathbf{D}$ | $\mathbf{A}$ | $\mathbf{C}$ | $\mathbf{B}$ |
| :---: | :---: | :---: | :---: |
| $\mathbf{B}$ | $\mathbf{C}$ | $\mathbf{A}$ | $\mathbf{D}$ |
| $\mathbf{C}$ | $\mathbf{D}$ | $\mathbf{B}$ | $\mathbf{A}$ |
| $\mathbf{A}$ | $\mathbf{B}$ | $\mathbf{D}$ | $\mathbf{C}$ |

## Latin Square Designs

- Factors:
- Row (blocking factor 1)
- Column (blocking factor 2)
- Treatment (factor of interest)
- Statistical Model
- Three-way ANOVA model
- (additive model with single replication)
- Everything is regression!
(Professor Scott Emerson)


