Lecture 6 – ANOVA and Linear Models

STAT/BIOF/GSAT 540: Statistical Methods for High Dimensional Biology

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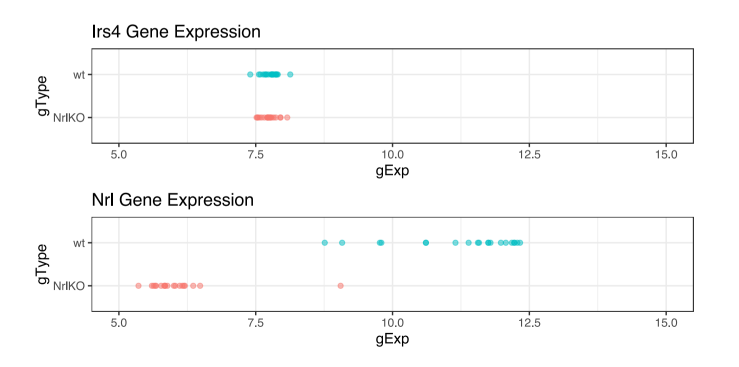
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Are these genes truly different in NrIKO compared to WT?

H₀: the expression level of gene A is the same in both conditions.

Is there **enough** evidence in the data to reject H₀?



Statistics: use a random sample to learn about the population

Population (unknown)

$$Y \sim F$$

 $Z \sim G$

$$E[Y] = \mu_Y$$

$$E[Z] = \mu_Z$$

Sample

(observed with randomness)

$$Y_1, Y_2, \ldots, Y_{n_Y}$$

$$Z_1, Z_2, \ldots, Z_{n_Z}$$

$$\hat{\mu}_Y = \bar{Y} = \frac{\sum_{i=1}^{n_Y} Y_i}{n_Y}$$

$$\hat{\mu}_Z = \bar{Z} = \frac{\sum_{i=1}^{n_Z} Z_i}{n_Z}$$

Last class: hypothesis testing

- 1. Define a test statistic to test H_0
 - 2-sample *t*-test
 - Welch *t*-test
 - Wilcoxon rank-sum test
 - Kolmogorov-Smirnov test
- 2. Compute the observed value for the test statistic
- 3. Compute the probability of seeing a test statistic as extreme as that observed, under the null sampling distribution (p-value)
- 4. Make a decision about the significance of the results, based on a pre-specified value (alpha, significance level)

We can run these tests in R

Example: use the t.test function to test H_0 using a classical 2-sample \emph{t} -test.

```
miniDat %>% subset(gene == "Irs4") %>% t.test(gExp ~ gType, data = .,
    var.equal = TRUE)
##
##
      Two Sample t-test
##
## data: gExp by gType
## t = -0.52865, df = 37, p-value = 0.6002
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.12597002 0.07383844
## sample estimates:
## mean in group NrlKO mean in group wt
##
             7.739684
                                 7,765750
```

Today...

- show how to compare means of different groups (2 or more) using a linear regression model
 - dummy variables to model the levels of a qualitative explanatory variable
- write a linear model using matrix notation
 - understand which matrix is built by R
- distinguish between conditional and marginal effects
 - *t*-tests vs *F*-tests

```
> t.test(gExp ~ gType, miniDat,
+ subset = gene == "Irs4", var.equal = TRUE)
```

two sample t test

```
H_0: \mu_1 = \mu_2
```

```
> summary(aov(gExp ~ gType, miniDat,
+ subset = gene == "Irs4"))
```

(one-way) analysis of variance "ANOVA"

```
> summary(lm(gExp ~ gType, miniDat,
+ subset = gene == "Irs4"))
```

linear model linear regression

It seems that we can use any of these methods to test H_0

```
> t.test(gExp ~ gType, miniDat,
         subset = gene == "Irs4", var.equal = TRUE)
    Two Sample t-test
                                                    Irs4 gene
data: qExp by qType
t = 0.5286, df = 37, p-value = 0.6002
                                                              mean = 7.74
                                               NrlKO -
<snip, snip>
sample estimates:
   mean in group wt mean in group NrlKO
                                                                mean = 7.77
           7.765750
                                7.739684
                                                 wt -
> summary(aov(gExp ~ gType, miniDat,
           subset = gene == "Irs4"))
                                                            7.6
                                                                    7.8
                                                    7.4
                                                                            8.0
            Df Sum Sq Mean Sq F value Pr(>F)
                                                                 gExp
gType 1 0.0066 0.00662 0.279
                                        0.6
                                                     7.739684 - 7.765750 = -0.026066
Residuals 37 0.8764 0.02369
                                                      -0.5286494 ^ 2 = 0.2794702
```

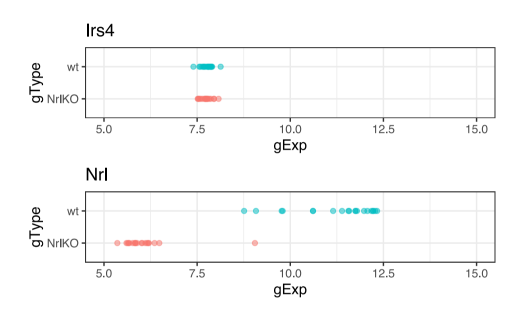
+ subset = gene == "Irs4"))
<snip, snip>
Coefficients:

> summary(lm(qExp ~ qType, miniDat,

t-test *vs* linear regression: why the same results?

```
irs4Dat <- subset(miniDat,gene=="Irs4")</pre>
ttest.irs4<-t.test(gExp ~ gType, irs4Dat, var.equal = TRUE)</pre>
list("t value"=ttest.irs4$stat,"p-value"=ttest.irs4$p.value)
## $t value
##
## -0.5286494
##
## $p-value
## [1] 0.6002058
lm.irs4 <- summary(lm(gExp ~ gType, irs4Dat))</pre>
list("t value"=lm.irs4$coeff[2,3],"p-value"=lm.irs4$coeff[2,4])
## $t value
## [1] 0.5286494
##
## $p-value
## [1] 0.6002058
```

t-test *vs* linear regression: where's the line?



Note that the *y*-axis in these plots is not numerical, thus a line in this space does not have any mathematical meaning.

Why can we run a t-test with a linear regression model?

From *t*-test to linear regression

Let's change the notation to give a common framework to all methods

$$Y\sim G;\; E[Y]=\mu_Y$$
 \downarrow $Y=\mu_Y+arepsilon_Y;\; arepsilon_Y\sim G;\; E[arepsilon_Y]=0$

We can use a subindeces to distinguish observations from each group, i.e.,

$$Y_{ij} = \mu_j + arepsilon_{ij}; \;\; arepsilon_{ij} \sim G_j; \;\; E[arepsilon_{ij}] = 0;$$

where $j = \{\text{wt}, \text{NrlKO}\}\$ or $j = \{1, 2\}$ identifies the groups; and $i = 1, \dots, n_j$ identifies the observations within each group

For example: Y_{11} is the first observation in group 1 or WT

The goal is to test

$$H_0:\mu_1=\mu_2$$

using data from the model

$$Y_{ij} = \mu_j + arepsilon_{ij}; \;\; arepsilon_{ij} \sim G; \;\; E[arepsilon_{ij}] = 0;$$

where $j = \{ \text{wt}, \text{NrlKO} \}$ or $j = \{1, 2\}$; and $i = 1, \ldots, n_j$.

For simplicity, we assume a common distribution G for all groups

Note that the population means are given by $E[Y_{ij}]=\mu_j$, i.e., the model is written with a cell-means - μ_j - parametrization

Note that for each group, the population mean is given by

$$E[Y_{ij}] = \mu_j,$$

A natural **estimator** of the population mean is the **sample mean**

Classical hypothesis testing methods use the group sample means as estimators

7,765750

See, for example, the t.test function in R:

7,739684

##

```
ttest.irs4$estimate

## mean in group NrlKO mean in group wt
```

However, the \tag{Tm function reports other estimates, why?

```
(means.irs4 <- as.data.frame(irs4Dat %>% group_by(gType) %>%
    summarize(meanGroups = mean(gExp, digits = 6))))
    gType meanGroups
## 1 NrlKO 7.739684
## 2
       wt 7,765750
lm.irs4$coefficients[,1]
## (Intercept) gTypewt
## 7.73968421 0.02606579
(Intercept) is the sample mean of NrlKO group
                                                  but gTypewt is not the sample mean of the WT
                                                  group
```

Parametrizations: which parameters we use to write the model?

By default, the lm does not use the cell-means parametrization The goal is to *compare* the means, not to study each in isolation

From **cell-means** - μ_j :

$$Y_{ij} = \mu_j + arepsilon_{ij}; \;\; arepsilon_{ij} \sim G; \;\; E[arepsilon_{ij}] = 0;$$

to reference-treatment effect - (θ, τ_j) :

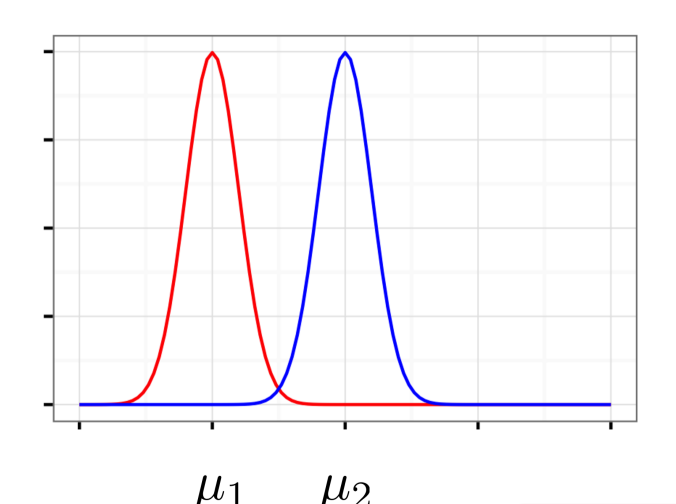
$$Y_{ij}= heta+ au_j+arepsilon_{ij}; \;\; au_1=0, \;\; arepsilon_{ij}\sim G; \;\; E[arepsilon_{ij}]=0;$$

Note that for each group, the population mean is given by

$$E[Y_{ij}] = heta + au_j = \mu_j,$$

and $au_2=\mu_2-\mu_1=E[Y_{i2}]-E[Y_{i1}]$ compares the means

Relation between parametrizations



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Im reports the sample mean of the reference group (NrIKO): $\hat{ heta}$

and the treatment effect, i.e., difference between the sample means of both groups: $\hat{ au}_2$

```
lm.irs4$coefficients[, 1]

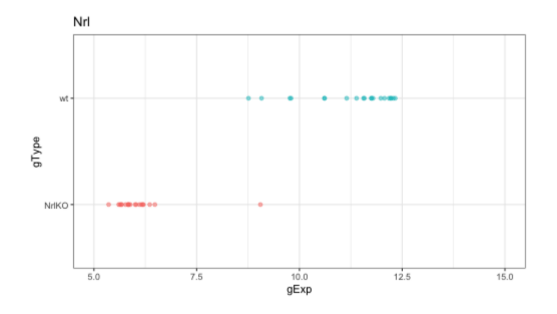
## (Intercept) gTypewt
## 7.73968421 0.02606579

data.frame(meanWT = means.irs4[1, 2],
    meanDiff = diff(means.irs4$meanGroups))

## meanWT meanDiff
## 1 7.739684 0.02606579
```

We still haven't answered ... where's the line??

$$Y_{ij}= heta+ au_j+arepsilon_{ij}; \;\; au_1=0, \;\; arepsilon_{ij}\sim G; \;\; E[arepsilon_{ij}]=0;$$



Dummy variables

Let's re-write our model using dummy (or indicator) variables:

$$Y_{ij} = heta + au_j + arepsilon_{ij}; \;\; au_1 = 0, \;\; arepsilon_{ij} \sim G; \;\; E[arepsilon_{ij}] = 0;
onumber \ Y_{ij} = heta + au_2 imes x_{ij} + arepsilon_{ij}; \;\; x_{ij} = \left\{ egin{array}{l} 1 ext{ if } j = 2 \ 0 ext{ otherwise} \end{array}
ight.$$

Note that $Y_{i1}= heta+arepsilon_{i1}$, because $au_1=0$ and $x_{i1}=0$ and $Y_{i2}= heta+ au_2+arepsilon_{i2}$, because $x_{i2}=1$ (for all i)

The second form is written as a linear ($y=a+bx+\varepsilon$) regression, with a special (dummy) explanatory variable x_{ij}

Using a dummy variables to model our categorical variables gtype we can perform a 2-sample *t*-test with a linear model

$$Y_{ij} = heta + au_2 imes x_{ij} + arepsilon_{ij}; \;\; x_{ij} = \left\{egin{array}{l} 1 ext{ if } j = 2 \ 0 ext{ if } j = 1 \end{array}
ight.$$

[1] 0.5286494

[1] 0.6002058

\$p-value

##

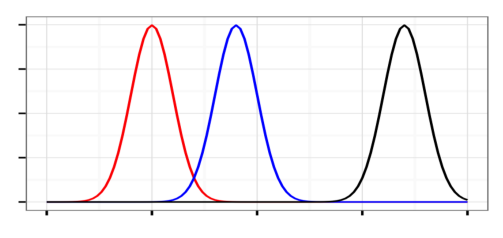
Beyond 2-groups comparisons: difference of means

"cell-means"

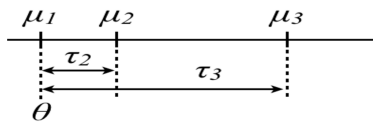
$$Y_{ij} = \mu_j + \varepsilon_{ij}$$

"reference-treatments"

$$Y_{ij} = \theta + \tau_j + \varepsilon_{ij}, (\tau_1 = 0)$$



More than 2 groups!



Dummy variables can be used to model one *or more* categorical variables with 2 *or more* levels!

2-sample *t*-test using a linear model

$$Y_{ij} = heta + au_2 imes x_{ij} + arepsilon_{ij}; \;\; x_{ij} = \left\{ egin{array}{l} 1 ext{ if } j = 2 \ 0 ext{ if } j = 1 \end{array}
ight.$$

1-way ANOVA with many levels (*) using a linear model

$$Y_{ij} = heta + au_2 imes x_{ij2} + au_3 imes x_{ij3} + arepsilon_{ij}; \;\; x_{ij2} = \left\{egin{array}{l} 1 ext{ if } j = 2 \ 0 ext{ otherwise} \end{array}; \; x_{ij3} = \left\{egin{array}{l} 0 ext{ if } j = 3 \ 1 ext{ otherwise} \end{array}
ight.$$

This is why R can estimate all of them with lm()

(*) in general, *yet* another parametrization is used to present ANOVA

t-test

Special case of ANOVA, but with ANOVA you can compare **more than two groups** and **more than one factor**.

ANOVA

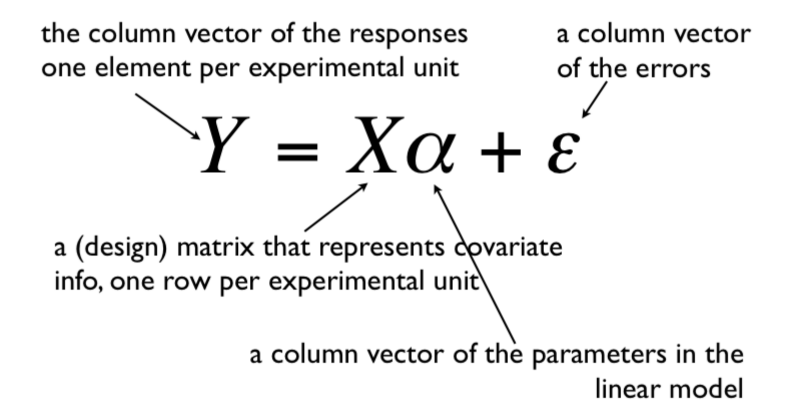
Special case of linear regression, but with linear regression you can include **quantitative** variables in the model.

Linear regression

Provides a unifying framework to model the association between a response **many quantitative** and qualitative variables.

In R: all can be computed using the lm () function.

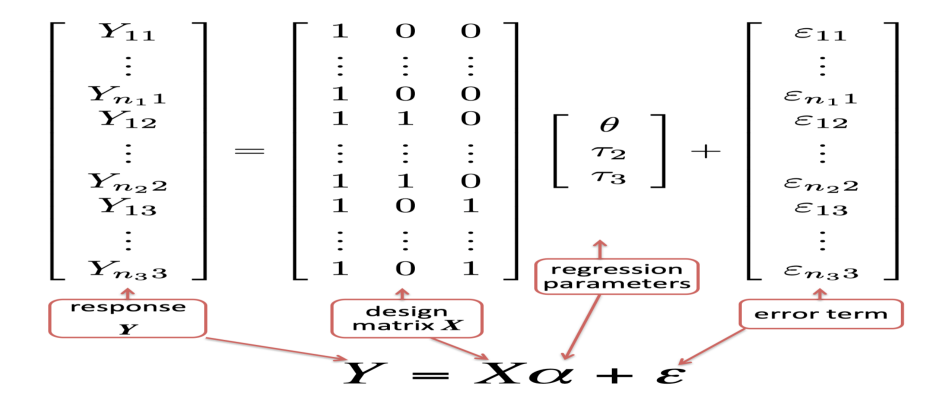
Linear models using matrix notation



It will become handy to write our model using matrix notation

Let's form an X matrix for a 3-groups comparison:

$$Y_{ij} = heta + au_2 imes x_{ij2} + au_3 imes x_{ij3} + arepsilon_{ij}$$



Note that x_{ij2} and x_{ij3} become the 2nd and 3rd columns of X: $x_{i12} = x_{i13} = 0$ for the reference group; $x_{i22} = 1$ for the 2nd group; and $x_{i33} = 1$ for the 3rd group

$$Y_{ij} = \theta + \tau_2 \times x_{ij2} + \tau_3 \times x_{ij3} + \varepsilon_{ij}$$

$$\begin{bmatrix} Y_{11} \\ \vdots \\ Y_{n_{1}1} \\ Y_{12} \\ \vdots \\ Y_{n_{2}2} \\ Y_{13} \\ \vdots \\ Y_{n_{3}3} \end{bmatrix} = \begin{bmatrix} 1 & 0 & 0 \\ \vdots & \vdots & \vdots \\ 1 & 0 & 0 \\ 1 & 1 & 0 \\ \vdots & \vdots & \vdots \\ 1 & 1 & 0 \\ 1 & 0 & 1 \end{bmatrix} \begin{bmatrix} \theta \\ \tau_{2} \\ \tau_{3} \end{bmatrix} \end{bmatrix} + \begin{bmatrix} \varepsilon_{11} \\ \vdots \\ \varepsilon_{n_{1}1} \\ \varepsilon_{12} \\ \vdots \\ \varepsilon_{n_{2}2} \\ \varepsilon_{13} \\ \vdots \\ \varepsilon_{n_{3}3} \end{bmatrix}$$

Note that
$$Y_{i1}=1 imes heta+0 imes au_2+0 imes au_3+arepsilon_{i1}= heta+arepsilon_{i1}$$

Note that $Y_{i2}=1 imes heta+1 imes au_2+0 imes au_3+arepsilon_{i2}= heta+ au_2+arepsilon_{i2}$
Note that $Y_{i3}=1 imes heta+0 imes au_2+1 imes au_3+arepsilon_{i3}= heta+ au_3+arepsilon_{i3}$

Which is the same as $\;
ightarrow\; Y_{ij}= heta+ au_j+arepsilon_{ij},\; au_1=0$

Which is the same as $\ o \ Y_{ij} = heta + au_2 imes x_{ij2} + au_3 imes x_{ij3} + arepsilon_{ij}$

$$Y=Xlpha+\mathcal{E}$$

$$\begin{bmatrix} Y_{11} \\ Y_{21} \\ \vdots \\ Y_{n_33} \end{bmatrix} = \begin{bmatrix} 1 & 0 & 0 \\ \vdots & \vdots & \vdots \\ 1 & 0 & 0 \\ 1 & 1 & 0 \\ \vdots & \vdots & \vdots \\ 1 & 1 & 0 \\ 1 & 0 & 1 \end{bmatrix} \begin{bmatrix} \theta \\ au_2 \\ au_3 \end{bmatrix} + \begin{bmatrix} arepsilon_{11} \\ arepsilon_{21} \\ \vdots \\ arepsilon_{n_33} \end{bmatrix}$$

$$\mu_2-\mu_1$$

Note that the model is still written with a reference-treatment parametrization (difference of means)

$$egin{align} E[Y_{i1}] &= heta \ & E[Y_{i2}] = heta + au_2 \ o au_2 = E[Y_{i2}] - E[Y_{i1}] = \mu_2 - \mu_1 \ & E[Y_{i3}] = heta + au_3 \ o au_3 = E[Y_{i3}] - E[Y_{i1}] = \mu_3 - \mu_1 \ \end{aligned}$$

Linear regression can include quantitative & qualitative covariates.

$$Y = X\alpha + \varepsilon$$

This gives us a VERY FLEXIBLE framework!!

$$\begin{bmatrix} 1 & 0 & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots \\ 1 & 0 & 0 & 0 \\ 1 & 1 & 0 & 0 \\ 1 & 1 & 0 & 0 \\ \vdots & \vdots & \vdots & \vdots \\ 1 & 1 & 0 & 0 \\ 1 & 0 & 1 & 0 \\ \vdots & \vdots & \vdots & \vdots \\ 1 & 0 & 1 & 0 \\ 1 & 0 & 0 & 1 \\ \vdots & \vdots & \vdots & \vdots \\ 1 & 1 & 1 & 1 \\ \vdots & \vdots & \vdots & \vdots \\ 1 & 1 & 1 & 1 \\ \vdots & \vdots & \vdots & \vdots \\ 1 & 1 & 1 & 1 \\ \vdots & \vdots & \vdots & \vdots \\ 1 & 1 & 1 & 1 \\ \vdots & \vdots & \vdots & \vdots \\ 1 & 1 & 1 & 1 \\ \vdots & \vdots & \vdots & \vdots \\ 1 & 1 & 1 & 1 \end{bmatrix}$$

$$\begin{bmatrix} 1 & 0 & 1.22 & 0 \\ 1 & 0 & 2.02 & 0 \\ 1 & 0 & 1.42 & 0 \\ \vdots & \vdots & \vdots & \vdots \\ 1 & 0 & 1.89 & 0 \\ 1 & 1 & 2.01 & 2.01 \\ \vdots & \vdots & \vdots & \vdots \\ 1 & 1 & 1.56 & 1.56 \\ 1 & 1 & 2.17 & 2.17 \\ 1 & 1 & 1.51 & 1.51 \end{bmatrix}$$

1 categorical covariate

2 categorical covariates 1 continuous covariate

1 continuous 1 categorical

AND MANY MORE

Tip: ?model.matrix

Linear in the parameters α : X can contain x^2 , log(x), etc.

How it works in practice using Im() in R

$$Y = X\alpha + \varepsilon$$

 $lm(y \sim x, data = yourData)$

y ~ x: formula, y numeric, x numeric and/or factor yourData: data.frame in which x and y are to be found (optional but recommended)

By default, R uses a ref-tx parametrization but you can control that!

$$Y = X\alpha + \varepsilon$$

- Mathematically, \boldsymbol{X} is a numeric matrix
- If your data contains categorical variables (e.g., gType), you need to set them as **factors**
- R creates appropriate dummy variables for factors!

```
str(irs4Dat$gType)
```

Factor w/ 2 levels "NrlKO","wt": 2 2 2 2 1 1 1 2 2 2 ...

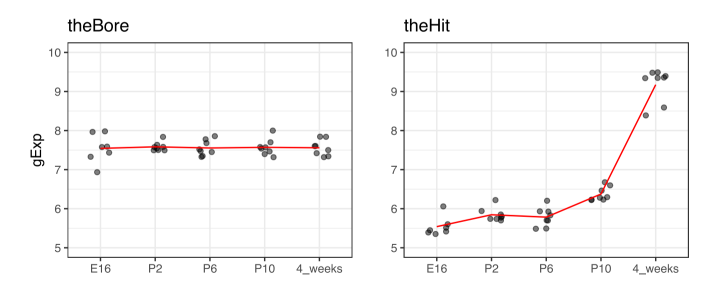
Under the hood, R creates a numeric X:

```
data.frame(X = model.matrix(gExp ~ gType, irs4Dat),
     gType = irs4Dat$gType) %>% head(10)
##
     X..Intercept. X.gTypewt gType
## 1
                                wt
## 2
                               wt
## 3
                                wt
## 4
                                 wt
## 5
                            0 NrlKO
## 6
                            0 NrlKO
## 7
                            0 NrlKO
## 8
                                 wt
## 9
                                wt
## 10
                                 wt
```

Beyond 2-group comparisons in our case study:

Is the expression of gene A the same at all developmental stages?

$$H_0: \mu_{E16} = \mu_{P2} = \mu_{P6} = \mu_{P10} = \mu_{4W}$$

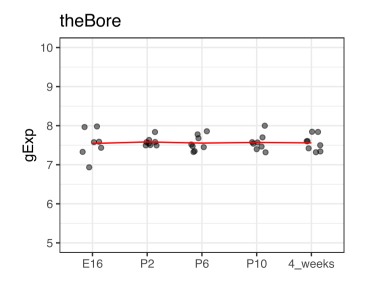


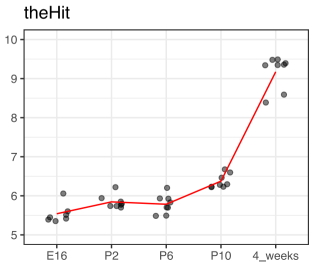
Note: 4W = 4_weeks

The sample means: $\hat{\mu}_{E16},~\hat{\mu}_{P2},~\hat{\mu}_{P6},~\hat{\mu}_{P10},~\hat{\mu}_{4W}$

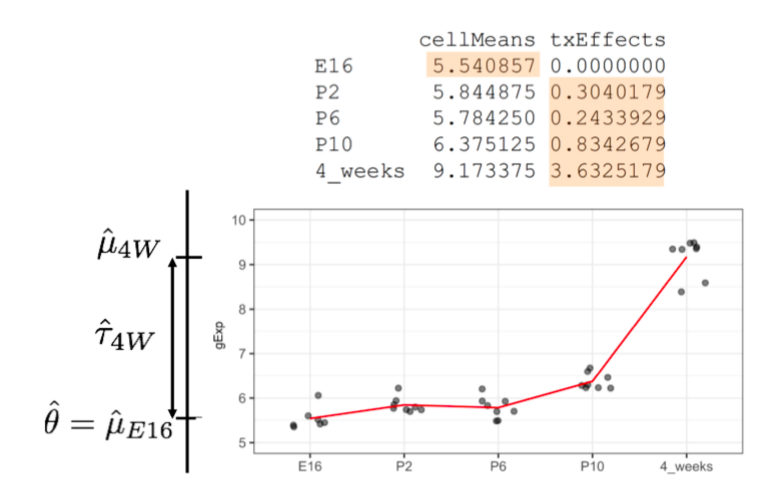
```
with(devDat, tapply(gExp, list(devStage, gene), mean))
```

```
## E16 7.544143 5.540857
## P2 7.583500 5.844875
## P6 7.554000 5.784250
## P10 7.571000 6.375125
## 4_weeks 7.559000 9.173375
```





"theHit" with significant time ("treatment") effect



"theHit" with significant time ("treatment") effect

Can you guess the size of the X matrix??

How many dummy variables do we need?

"theHit" with significant time ("treatment") effect

We need 4 dummy variables to estimate and test 4 time differences:

```
x_{P2}: P2 vs E16, x_{P6}: P6 vs E16, x_{P10}: P10 vs E16, x_{4W}: 4W vs E16)
```

Mathematically:

$$Y_{ij} = heta + au_{P2} imes x_{ijP2} + au_{P6} imes x_{ijP6} + au_{P10} imes x_{ijP10} + au_{4W} imes x_{ij4W} + arepsilon_{ij}$$

Notation: x_{ijk} , where i is an index for the observation, j for the level of devStage, and k for the name of the dummy variable

Under the hood, R creates a numeric X:

```
X.matrix <- data.frame(X = model.matrix(gExp ~ devStage, irs4Dat),
    devStage = irs4Dat$devStage)</pre>
```

```
##
      X..Intercept. X.dStP2 X.dStP6 X.dStP10 X.dS4W
## 1
                                                    0 E16
                           0
## 2
                                                    0 E16
                           0
## 3
                                                    0 E16
                           0
## 4
                                                    0 E16
                           0
## 5
                                                    0 E16
                           0
## 6
                                                    0 E16
                           0
## 7
                                                    0 E16
                           0
## 8
                                                      P2
## 9
                                                    0 P2
## 10
                                                    0 P2
## 11
                                                      P2
## 12
                                                       P2
## 13
                                                    0 P2
## 14
                                                       P2
## 15
                                                       P2
## 16
                           0
                                                       P6
```

Note: column names changed and first 16 rows displayed to fit output in the page (code hidden)

```
summary(lm(gExp~devStage,subset(devDat,gene=="theHit")))$coeff
##
                   Estimate Std. Error t value
                                                   Pr(>|t|)
## (Intercept)
                  5.5408571 0.1021381 54.248698 1.307554e-34
## devStageP2
                 0.3040179 0.1398583 2.173756 3.678022e-02
## devStageP6 0.2433929 0.1398583 1.740282 9.085489e-02
## devStageP10
                 0.8342679 0.1398583 5.965093 9.559065e-07
## devStage4 weeks 3.6325179 0.1398583 25.972843 5.266481e-24
means.dev %>% mutate(txEffects=cellMeans-cellMeans[1])
    devStage cellMeans txEffects
##
## 1
         E16 5.540857 0.0000000
## 2
     P2 5.844875 0.3040179
## 3
    P6 5.784250 0.2433929
## 4
         P10 6.375125 0.8342679
## 5 4 weeks 9.173375 3.6325179
```

Estimate: $\hat{ heta}=\hat{\mu}_{E16}=ar{Y}_{.E16}$

 $H_0: heta=0$ or

 $H_0: \mu_{E16} = 0$

we are not usually interested in testing this hypothesis

```
summary(lm(gExp~devStage,subset(devDat,gene=="theHit")))$coeff
##
                   Estimate Std. Error
                                         t value
                                                     Pr(>|t|)
## (Intercept)
                  5.5408571 0.1021381 54.248698 1.307554e-34
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means.dev %>% mutate(txEffects=cellMeans-cellMeans[1])
     devStage cellMeans txEffects
##
## 1
         E16 5.540857 0.0000000
## 2
          P2 5.844875 0.3040179
## 3
          P6 5.784250 0.2433929
## 4
         P10 6.375125 0.8342679
## 5 4 weeks 9.173375 3.6325179
```

Estimate:

$$\hat{ au}_{P2} = \hat{\mu}_{P2} - \hat{\mu}_{E16} = ar{Y}_{.P2} - ar{Y}_{.E16}$$

 $H_0: au_{P2} = 0 ext{ or }$

 $H_0: \mu_{P2} = \mu_{E16}$

we *are* usually interested in testing this hypothesis: first 2 days after birth

```
summary(lm(gExp~devStage,subset(devDat,gene=="theHit")))$coeff
##
                   Estimate Std. Error t value
                                                   Pr(>|t|)
## (Intercept)
                  5.5408571 0.1021381 54.248698 1.307554e-34
## devStageP2
                  0.3040179 0.1398583 2.173756 3.678022e-02
## devStageP6 0.2433929 0.1398583 1.740282 9.085489e-02
## devStageP10
                  0.8342679 0.1398583 5.965093 9.559065e-07
## devStage4 weeks 3.6325179 0.1398583 25.972843 5.266481e-24
means.dev %>% mutate(txEffects=cellMeans-cellMeans[1])
    devStage cellMeans txEffects
##
## 1
         E16 5.540857 0.0000000
## 2
     P2 5.844875 0.3040179
## 3
    P6 5.784250 0.2433929
         P10 6.375125 0.8342679
## 4
```

Estimate:

$$\hat{ au}_{4W} = \hat{\mu}_{4W} - \hat{\mu}_{E16} = ar{Y}_{.4W} - ar{Y}_{.E16}$$

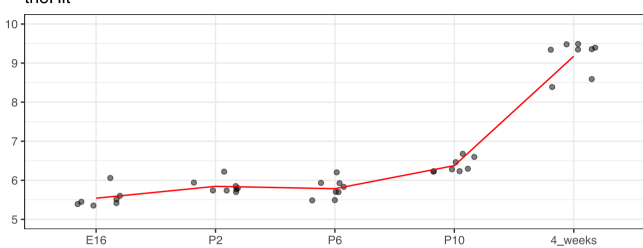
5 4 weeks 9.173375 3.6325179

 $H_0: au_{4W} = 0 ext{ or }$

 $H_0: \mu_{4W}=\mu_{E16}$

we *are* usually interested in testing this hypothesis: 4 weeks after birth





$$Y = Xlpha + arepsilon$$
 $lpha = (heta, au_{P2}, au_{P6}, au_{P10}, au_{4W})$

We generally test two types of null hypotheses:

$$H_0: au_i=0$$

VS

$$H_0: au_j
eq 0$$

for each *j* individually

e.g., Is gene A differencially expressed 2 days after birth?

$$H_0:\tau_{P2}=0$$

$$H_0: au_j=0$$

VS

$$H_0: au_j
eq 0$$

for all *j* at the same time

e.g., Is gene A significantly affected by time (devStage)?

$$H_0: au_{P2} = au_{P6} = au_{P10} = au_{4W} = 0$$

Two types of null hypotheses in R:

$$Y = X\alpha + \varepsilon$$
 $\alpha = (\theta, \tau_{P2}, \tau_{P6}, \tau_{P10}, \tau_{4 \text{ weeks}})$

```
H_0: \tau_j = 0
vs H_0: \tau_j \neq 0 for each j individually
```

```
H_0: 	au_j = 0 AND statement vs H_0: 	au_j \neq 0 OR statement for all j at the same time
```

```
> summary(hitFit)
Call:
lm(formula = gExp ~ devStage, <blah, blah>)
<snip, snip>
Coefficients:
               Estimate Std. Error t value Pr(>|t|)
                           0.1021 54.249 < 2e-16 ***
                 5.5409
(Intercept)
                0.3040
                           0.1399 2.174 0.0368 *
devStageP2
                0.2434 0.1399 1.740 0.0909 .
devStageP6
           0.8343
                           0.1399 5.965 9.56e-07 ***
devStageP10
                           0.1399 25.973 < 2e-16 ***
devStage4 weeks 3.6325
<snip, snip>
F-statistic: 243.4 on 4 and 34 DF, p-value: < 2.2e-16
```

F-test and overall significance of one or more covariates

• the *t*-test in linear regression allows us to test single hypotheses:

$$H_0: au_i=0$$

$$H_A: au_j
eq 0$$

• but we often like to test multiple hypotheses *simultaneously*:

$$H_0: au_{P2} = au_{P6} = au_{P10} = au_{4W} = 0 \ [ext{AND statement}]$$

$$H_A: au_i
eq 0 ext{ for some i [OR statement]}$$

the *F*-test allows us to test such compound tests

To conclude

- we can use different parametrizations to write statistical models

From **cell-means** -
$$\mu_j$$
: $Y_{ij}=\mu_j+arepsilon_{ij}; \ \ arepsilon_{ij}\sim G; \ \ E[arepsilon_{ij}]=0;$ to **reference-treatment effect** - $(heta, au_j)$: (used by default by lm) $Y_{ij}= heta+ au_j+arepsilon_{ij}; \ \ au_1=0, \ \ arepsilon_{ij}\sim G; \ \ E[arepsilon_{ij}]=0;$

- we can compare group means (2 or more) using a linear model
 - dummy variables (e.g., x_{ijP2}) to model the levels of a qualitative explanatory variables

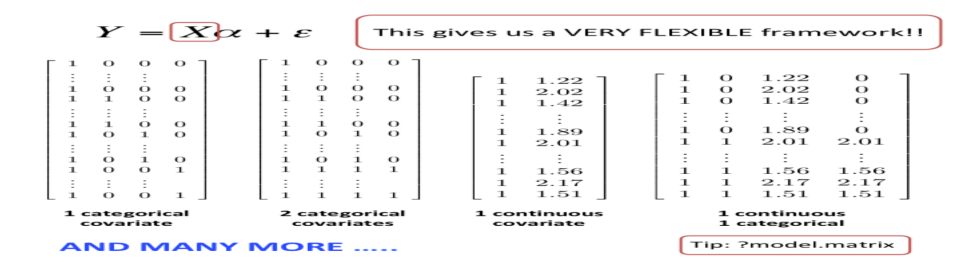
$$Y_{ij} = heta + au_{P2} imes x_{ijP2} + au_{P6} imes x_{ijP6} + au_{P10} imes x_{ijP10} + au_{4W} imes x_{ij4W} + arepsilon_{ij}$$

• qualitative variables need to be set as "factors" in the data --> R creates the dummy variables

- we can write a linear model using matrix notation:

$$Y = X\alpha + \varepsilon$$

- Linear model can include quantitative & qualitative covariates.



- distinguish between single and joint hypotheses:
 - t-tests vs F-tests