Linear model parameters

Objectives

- Understand how to interpret R output and parameters in linear models
- Be able to describe the difference between an interactive and additive model
- Plot predictions from additive and interactive linear models

Model parameters: definitions

- · Parameters of a linear model typically characterize *differences* in means
- These are differences per unit of change for continuous predictors,
- These are differences between groups (or between group averages) for categorical predictors
- Interactions are differences between differences

Coding for categorical predictors: contrasts

- What do the parameters of a linear model mean?
- Start with categorical variables, because they're potentially more confusing ("intercept and slope" isn't too hard)
- Default R behaviour: treatment contrasts
 - β_1 = expected value in baseline group (= first level of the factor variable, by default the first in alphabetical order);
 - β_i = expected difference between group i and the first group.

Example

The previously explored ant-colony example:

Define data:

pr(lm1 <- lm(colonies~place,data=ants))</pre>

```
## (Intercept) 10.75 0.98 10.97 4.2e-06
## placeforest -3.75 1.27 -2.96 0.018
```

- The (Intercept) row refers to β_1 , which is the mean density in the "field" sites ("field" comes before "forest").
- The placeforest row indicates we are looking at the effect of forest level of the place variable, i.e. the difference between "forest" and "field" sites. (To know that "field" is the baseline level we must (1) remember, or look at levels(ants\$place) or (2) notice which level is missing from the list of parameter estimates.)

R's behaviour may seem annoying at first – it seems like the estimated values of the groups are what we're really interested in – but it is really designed for testing *differences among groups*. To get the estimates per group, you could:

 use a regression formula colonies~place-1, or equivalently colonies~place+0, to suppress the implicit intercept term:

```
## Estimate Std. Error t value Pr(>|t|)
## placefield 10.75 0.98 10.97 4.2e-06
## placeforest 7.00 0.80 8.75 2.3e-05
```

When you use the colonies~place-1 formula, the meanings of the parameters change: β_1 is the same (mean of "field"), but β_2 is 'mean of "forest" rather than ("(field)-(forest)").

Interpretation using predict

Use the predict function:

2 7.00 5.154296 8.845704

Interpretation using effects package

Use the effects package:

```
library("effects")
summary(allEffects(lm1))
   model: colonies ~ place
##
##
## place effect
## place
## field forest
## 10.75 7.00
##
   Lower 95 Percent Confidence Limits
##
## place
## field forest
## 8.489484 5.154296
##
   Upper 95 Percent Confidence Limits
## place
      field forest
##
## 13.010516 8.845704
```

Interpretation using 1smeans package

Use the 1smeans package:

```
library("lsmeans")
lsmeans(lm1,specs=~place)

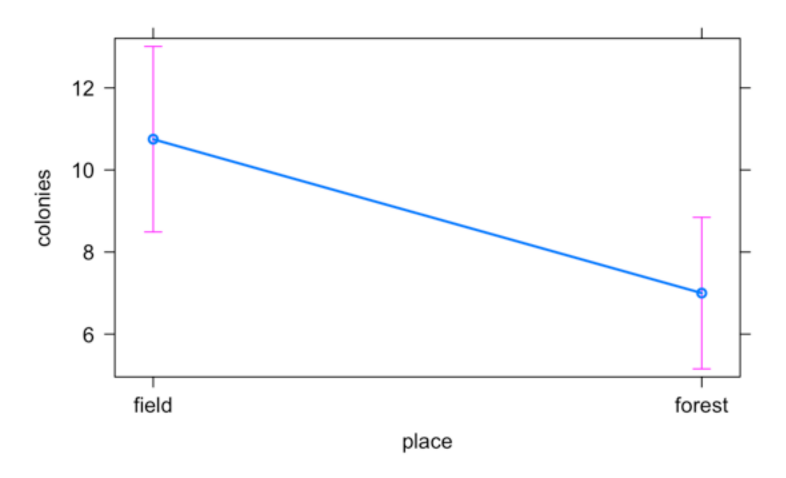
## place lsmean SE df lower.CL upper.CL
## field 10.8 0.98 8 8.49 13.01
## forest 7.0 0.80 8 5.15 8.85
##

## Confidence level used: 0.95
```

Graphical summaries from effects package

plot(allEffects(lm1))

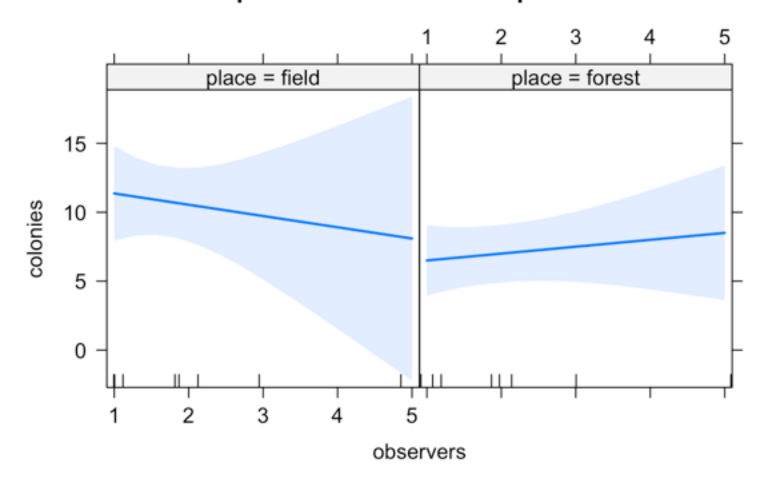
place effect plot



The effects package works on more complicated models

lm3 <- lm(colonies~place*observers,data=ants)
plot(allEffects(lm3))</pre>

place*observers effect plot

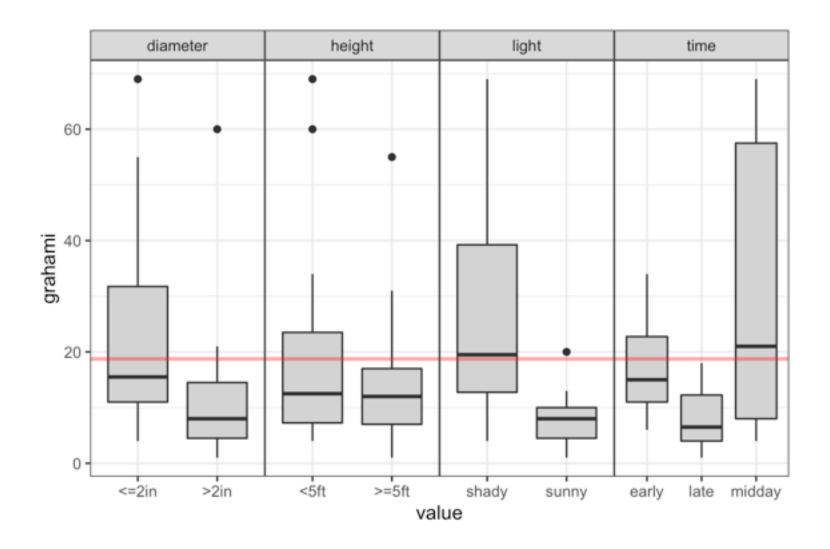


Switching to a dataset with more than two levels

Some data on lizard perching behaviour (brglm package; Schoener 1970 *Ecology* **51**:408-418).

```
lizards <- read.csv("lizards.csv")</pre>
```

Response is number of *Anolis grahami* lizards found on perches in particular conditions.



What is the effect of time of day on lizard perching?

```
pr(lmX <- lm(grahami~time,data=lizards))</pre>
```

```
## (Intercept) 17.63 5.97 2.95 0.0079
## timelate -9.50 8.44 -1.13 0.2739
## timemidday 14.52 8.74 1.66 0.1123
```

If we leave the factors alphabetical then β_1 ="early", β_2 ="late"-"early", β_3 ="midday"-"early". It might be more sensible to change the levels in accordance with time progression.

Change the order of the levels:

This just swaps the definitions of β_2 and β_3 .

```
pr(lmX <- lm(grahami~time,data=lizards))</pre>
```

```
## (Intercept) 17.63 5.97 2.95 0.0079
## timemidday 14.52 8.74 1.66 0.1123
## timelate -9.50 8.44 -1.13 0.2739
```

Multiple treatments and interactions

Additive model

Consider the light variable in addition to time.

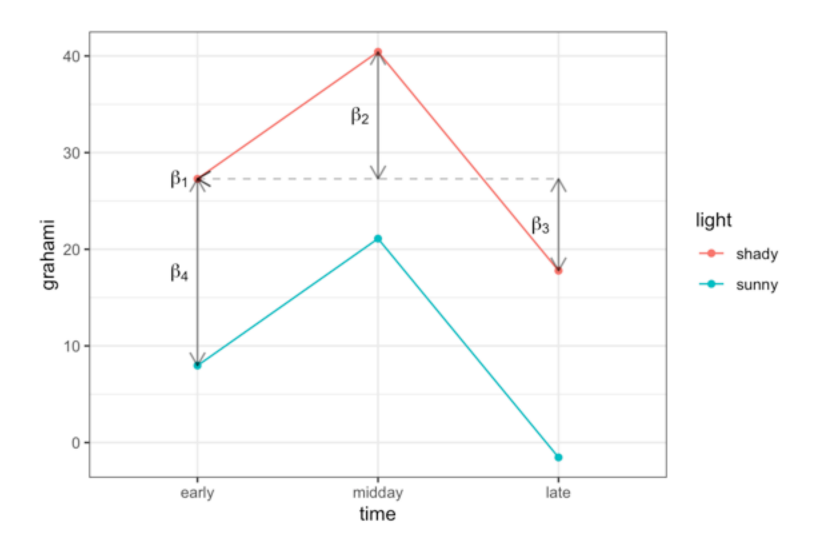
```
pr(lmTL1 <- lm(grahami~time+light,data=lizards))</pre>
```

```
## (Intercept) 27.29 5.63 4.85 0.00011
## timemidday 13.14 7.11 1.85 0.08010
## timelate -9.50 6.85 -1.39 0.18174
## lightsunny -19.32 5.73 -3.37 0.00321
```

- β_1 is the intercept ("early","shady");
- β_2 and β_3 are the differences from the baseline level ("early") of the *first* variable (time) in the *baseline* level of the other parameter(s) (light="shady");
- β_4 is the difference from the baseline level ("sunny") of the *second* variable (light) in the *baseline* level of time ("early").

Graphical interpretation

Loading required package: grid



What are the p-values?

```
pr(lmTL2 <- lm(grahami~time+light,data=lizards))</pre>
```

```
## (Intercept) 27.29 5.63 4.85 0.00011
## timemidday 13.14 7.11 1.85 0.08010
## timelate -9.50 6.85 -1.39 0.18174
## lightsunny -19.32 5.73 -3.37 0.00321
```

The p-values tell us the difference from the baseline level, not from each other

Assessing differences among pairs of variable levels

```
library(emmeans)
library(multcompView)
library(multcomp)
## Loading required package: mvtnorm
## Loading required package: survival
## Loading required package: TH.data
## Loading required package: MASS
##
## Attaching package: 'MASS'
## The following object is masked from 'package:dplyr':
##
      select
##
##
## Attaching package: 'TH.data'
                                                    21/33
```

Getting an ABCDEF.. list

```
lsm1<-emmeans(lmTL1,pairwise~time)</pre>
cld(lsm1$emmeans,by = NULL, Letters = "ABCDEFGHIJ")
   time
                   SE df lower.CL upper.CL .group
##
          emmean
##
   late 8.12 4.85 19
                            -2.02
                                     18.3
                                           A
   early 17.62 4.85 19 7.48
##
                                     27.8
                                           AB
   midday 30.76 5.20 19 19.89 41.6 B
##
##
## Results are averaged over the levels of: light
## Confidence level used: 0.95
## P value adjustment: tukey method for comparing a family o
## significance level used: alpha = 0.05
```

Interactions

- Interactions allow the value of one predictor to affect the relationship between another predictor and the response variable
- Interpreting main effects in the presence of interactions is tricky
- ullet Your estimate of the effect of variable B is no longer constant
- You need to pick a fixed point, or average in some way
- Example: $Y = a + b_1X_1 + b_2X_2 + b_{12}X_1 * X_2$
- The response to X_1 is $Y=(a+b_2X_2)+(b_1+b_{12}X_2)X_1$ the response to X_1 depends on the value of X_2 .

An example

- You think that the number of lizards on a perch on a sunny day might depend on the time day
- For example, on a very sunny day, there might be fewer lizards perching at noon; whereas on a cloudy day, the number of lizards might be highest at noon

Testing interactions

- Time is an important factor
- Use an interaction:

$$M = a + B_x X + B_t t + B_{xt} X t$$

- The interaction term $B_{\it xt}$ represents the difference in the response between the two groups.
- It asks: does the number of lizards perching depend on time of day and light?
- · Could also write:

$$M = a + B_1 light + B_2 time + B_3 light * time$$

Treatment and interactions

- We previously discussed the wrong construction of linear models to compare drug treatments in mutant and non-mutant mice (erroneous interactions)
- You want to know whether a drug that significantly reduces neuronal firing affects mutant or control mice differently
- You find that the drug sig. decreases neuronal firing in mutant mice
- The drug doesn't decrease sig. dec neuronal firing in non-mutants
- You need mouse_type*trmt to understand whether the treatment affects mice differently!

Interactions and parameters

- We can use CIs, and coefficient plots, and get a pretty good idea what's going on
- In more complicated cases, interaction terms may have many parameters
- These have all the interpretation problems of other multi-parameter variables
- Think about "differences in differences"

Interaction model

pr(lmTL2 <- lm(grahami~time*light,data=lizards))</pre>

```
Estimate Std. Error t value Pr(>|t|
##
                                     5.38
## (Intercept)
                                             4.37
                                                   0.0004
                          23.50
## timemidday
                                     7.60 3.65
                                                   0.0019
                          27.75
                                            -1.68
## timelate
                                                  0.1118
                         -12.75
                                     7.60
## lightsunny
                         -11.75
                                     7.60
                                            -1.55 0.1406
## timemidday:lightsunny -32.83
                                    11.19 -2.93
                                                  0.0092
                                    10.75 0.60
## timelate:lightsunny
                           6.50
                                                   0.5534
```

- Parameters β_1 to β_4 have the same meanings as before.
- β_5 and β_6 , labelled "timemidday:lightsunny" and "timelate:lightsunny" describe the difference between the expected mean value of these treatment combinations based on the additive model

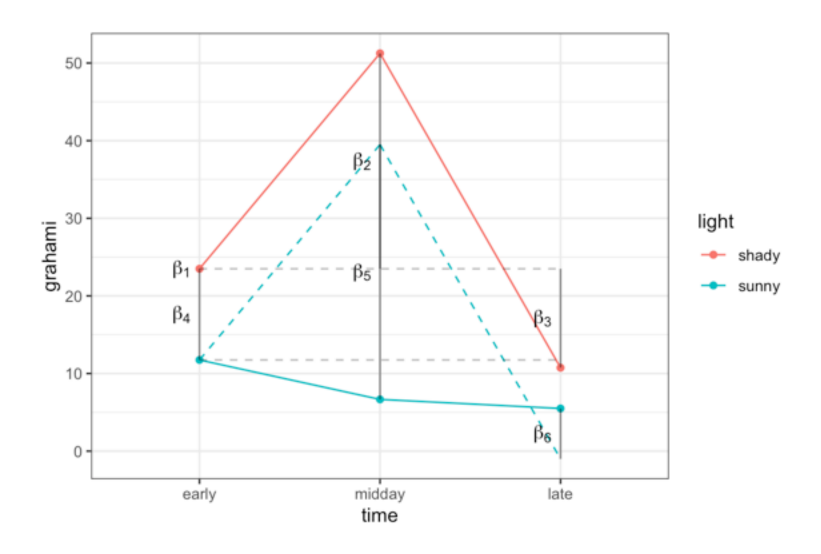
Interaction model cont

pr(lmTL2 <- lm(grahami~time*light,data=lizards))</pre>

```
##
                       Estimate Std. Error t value Pr(>|t|
## (Intercept)
                                     5.38
                                            4.37
                                                  0.0004
                          23.50
## timemidday
                                     7.60 3.65
                                                  0.0019
                          27.75
## timelate
                                     7.60
                                                 0.1118
                         -12.75
                                           -1.68
## lightsunny
                                           -1.55 0.1406
                         -11.75
                                    7.60
## timemidday:lightsunny -32.83 11.19 -2.93 0.0092
## timelate:lightsunny
                          6.50
                                    10.75 0.60
                                                  0.5534
```

- num lizards when sunny at midday = $\beta_1 + \beta_2 + \beta_4 + \beta_5$
- num lizards when sunny at late = $\beta_1 + \beta_3 + \beta_4 + \beta_6$

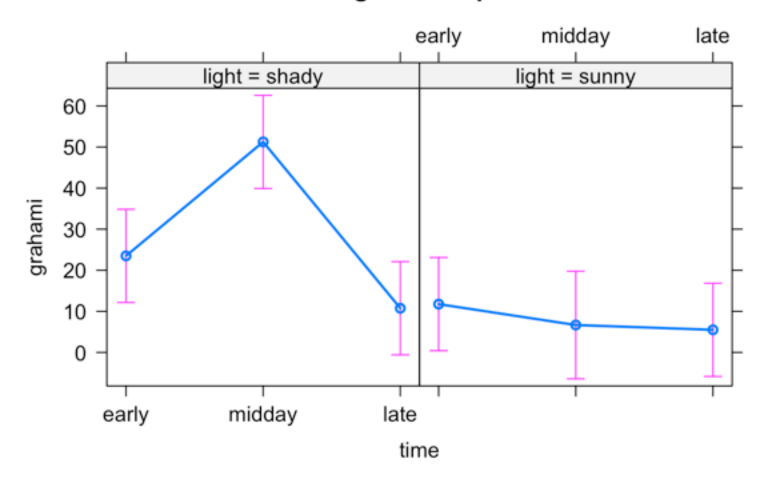
Graphical version



Effects plot

plot(allEffects(lmTL2))

time*light effect plot



Other refs

- http://sas-andr.blogspot.com/2010/10/example-89contrasts.html
- gmodels::fit.contrast() (show parameter estimates based on re-fitting models with new contrasts), rms::contrast.rms() (ditto, for rms-based fits)
- http://www.ats.ucla.edu/stat/r/library/contrast_codir

Assignment

- · PART 1
- 1. Make a univariate linear model for one of your hypotheses
- 2. Examine the assumptions of linearity (using tests or diagnostic plots)
- 3. Plot the relationship in ggplot using stat_smooth
- PART 2
- 1. Make a linear model (with more than one variable) for one of your hypotheses.

 Articulate which hypothesis you are testing.
- 2. Use an interactive model and an additive model. Explain what hypothesis each of these is testing, and what the R output is telling you about your data. (Hint: you can use Ismeans, effects, relevel, or predict to help you.) You should include this explanation in either your README or in your code.
- 3. Plot your model (e.g. using predict) and