Mixed-effects models in R

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Preface

This book is a collection of advanced topics in R. References to previous sections, as well as descriptions of datasets, are to the book by Duursma, Powell and Stone: *Data analysis and visualization with R*.

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Chapter 1

Mixed-effects models

In Chapter ??, we used linear models to estimate 'fixed' effects, which consist of specific and repeatable categories/variables that are representative of an entire population (e.g., species, age). In longitudinal studies (repeated measures) and in studies using hierarchical (nested) sampling, it is also possible to estimate effects associated with individuals sampled at random from the population of interest. These are 'random' effects and convey information about the degree that individuals in a population differ but not how or why they differ.

One way to differentiate fixed and random effects is that fixed effects contain levels that are informative beyond the current analysis (e.g., a species of tree or a specific management type) while random effects contain levels that are not informative beyond the current analysis (e.g., a group of trees within an observation plot or a field under specific management). Another way to understand the difference is that fixed effects influence the mean of the response while random effects influence the variance of the response.

Mixed-effects models estimate both fixed and random effects and are particularly useful when dealing with potential pseudoreplication and unbalanced designs. Including random effects can also account for variation that could mask patterns in an analysis considering only fixed effects.

Two commonly used packages for fitting mixed-effects models are nlme and lme4. In this chapter, we will use the newer lme4 package, but note that more complex correlation structures are only possible with the nlme package.

Rather than to present the theory underlying mixed-effects models, which is very complex, we will treat this topic by example, and thus aim at a practical application.

1.0.1 A note about p-values

The 1me4 package does not report p-values. The developers made this decision because p-values require calculating degrees of freedom. Random effects don't necessarily have to expend the same degrees of freedom as treating them as fixed effects, so the package developers have decided not to fudge this by calculating them for you.

You do have options though. See the help page ?pvalues for references to several options to calculate p-values with lme4.

In this chapter, we use two approaches to calculate p-values of fixed effects. To obtain the p-values of all fixed effects in a model, we use the Anova function from the car package (note that this function is distinct from anova!), but note that there are alternatives (and even when using Anova, you have alternatives).

The second approach is to use likelihood-ratio tests to test the significance of a single fixed effect in a model. To do this, we fit two models (one with, and one without the fixed effect of interest), and use anova on the two models to calculate a p-value for the fixed effect.

We recommend you try multiple methods to make sure that the main conclusions you draw from the results are robust.

1.1 A simple equation for a mixed-effects model

Before we fit a mixed-effects model, let's write down an equation that illustrates the roles of the fixed effects and random effects in the model.

For a simple linear regression of Y versus X (where X is numeric), we fit the model:

$$Y = \beta_0 + \beta_1 * X$$

where the β 's are the intercept and the slope of the fitted regression line.

A mixed-effects model additionally fits two random parameters, and we can write the model as:

$$Y = (\beta_0 + b_0) + (\beta_1 + b_1) * X$$

where b_0 is the random intercept (which is normally distributed with mean zero, and some standard deviation), and b_1 is the random slope (also assumed to be normally distributed with mean of zero and some standard deviation).

When we fit a mixed-effects model, not only do we get an estimate of the variance (or standard deviation) of the random effects, we also get estimates of this random effect for each individual. These estimates are known as the BLUPs (best linear unbiased predictors).

1.2 Example: individual-level variation

The following example uses the pref data (not yet described in Appendix A, download the file 'prefdata.csv'). The dataset contains measurements of leaf mass per area (LMA), and distance from the top of the tree (dfromtop) on 35 trees of two species. We want to know whether LMA decreases with dfromtop, as expected, and whether this decrease in LMA with distance from top differs by species.

```
# Read the data and inspect the first few rows, and the species factor variable.
pref <- read.csv("prefdata.csv")</pre>
head(pref)
                 species dfromtop totheight height
                             8.88
                                      22.40 13.52 319.4472 2.779190
## 1 FP11 Pinus ponderosa
## 2 FP11 Pinus ponderosa
                             0.62
                                      22.40 21.78 342.7948 4.010700
## 3 FP11 Pinus ponderosa
                           4.72
                                      22.40 17.68 329.5399 3.365579
                             2.74
5.48
                                      27.69 24.95 312.4467 3.682907
## 4 FP15 Pinus ponderosa
## 5 FP15 Pinus ponderosa
                                      27.69 22.21 278.4037 2.524224
## 6 FP15 Pinus ponderosa
                             8.40
                                      27.69 19.29 255.9716 2.351546
levels(pref$species)
## [1] "Pinus monticola" "Pinus ponderosa"
```

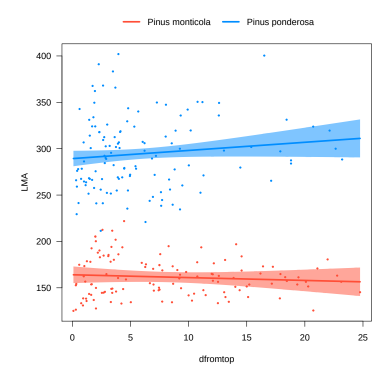


Figure 1.1: Leaf mass per area as a function of tree species (two colours) and the distance from the top of each tree, as fitted with a simple linear model and visualized with visreg.

Before we fit mixed-effects models, let's start with a linear regression that includes dfromtop and species as the predictor variables to observe the general patterns. We use visreg to quickly visualize the fitted linear model. The following code produces Fig. 1.1.

```
# Fit a linear regression by species (ignoring individual-level variation)
lm1 <- lm(LMA ~ species + dfromtop + species:dfromtop, data=pref)

# Plot predictions
library(visreg)
visreg(lm1, "dfromtop", by="species", overlay=TRUE)</pre>
```

As we can see in Fig. 1.1, there is a strong effect of species, but it appears that LMA and dfromtop are not significantly correlated.

Try this yourself Look at the anova table for the fitted linear regression model from the example above to confirm that LMA does not significantly change with dfromtop.

To see whether the relationship between LMA and dfromtop potentially varies from tree to tree, we fit a linear regression separately for each tree using the lmList function in the lme4 package and then plot the outcome (in Fig. 1.2).

```
# For the lmList function (Note: the nlme package also includes the lmList function)
library(lme4)

# fit linear regression by tree ('ID')
lmlis1 <- lmList(LMA ~ dfromtop | ID, data=pref)</pre>
```

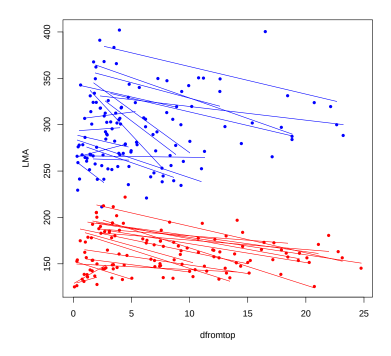


Figure 1.2: Leaf mass per area as a function of tree species (two colours) and the distance from the top of each tree. The solid lines represent the slope of the relationship for each individual tree.

```
# Extract coefficients (intercepts and slopes) for each tree
liscoef <- coef(lmlis1)</pre>
# load plottix for the 'ablineclip' function, which clips lines within the range of x
library(plotrix)
# split pref by tree (prefsp is a list)
prefsp <- split(pref, pref$ID)</pre>
# Plot
palette(c("red","blue"))
with(pref, plot(dfromtop, LMA, col=species, pch=16, cex=0.8))
for(i in 1:length(prefsp)){
  # Find min and max values of dfromtop, to send to ablineclip
  xmin <- min(prefsp[[i]]$dfromtop)</pre>
  xmax <- max(prefsp[[i]]$dfromtop)</pre>
  # add regression lines
  ablineclip(liscoef[i,1], liscoef[i,2], x1=xmin, x2=xmax,
             col=prefsp[[i]]$species)
}
```

From the figure we can conclude (informally) that:

1. Intercepts vary a lot between trees

- 2. There seems to be a negative relationship for many trees
- 3. It seems there is less variation between slopes than intercepts

We know the individual data points are not independent, as they are nested within trees (that is, multiple samples were collected for each tree). To properly account for this non-independence, we have to use a mixed-effects model. In the example below, we will fit two models: one with a random intercept only, and one with a random intercept and slope.

To specify random effects with 1mer, we add it to the formula in the right-hand side. For example, a random intercept for 'ID' (that is, the intercept will vary randomly among ID's) is coded as (1|ID). If we also allow the slope of the relationship to vary, we specify it as (dfromtop|ID) so that the slope and intercept of the relationship between LMA and dfromtop will vary randomly between tree ID's.

```
# Random intercept only
lme1 <- lmer(LMA ~ species + dfromtop + species:dfromtop + (1 | ID), data=pref)</pre>
# Random intercept and slope
lme2 <- lmer(LMA ~ species + dfromtop + species:dfromtop + (dfromtop | ID), data=pref)</pre>
# The AIC and a likelihood-ratio test tell us that we don't need a random slope.
# lower AIC indicates that model fit is better (more efficient)
AIC(lme1, lme2)
        df
                ATC
## lme1 6 2251.997
## lme2 8 2255.735
# Likelihood ratio test : the more complex model is not supported by the data.
# Note: the models will be re-fitted with ML instead of REML; this is necessary
# when performing likelihood-ratio tests.
anova(lme1, lme2)
## Data: pref
## Models:
## lme1: LMA ~ species + dfromtop + species:dfromtop + (1 | ID)
## lme2: LMA ~ species + dfromtop + species:dfromtop + (dfromtop | ID)
           AIC
                   BIC logLik deviance Chisq Chi Df Pr(>Chisq)
## lme1 6 2263.2 2284.3 -1125.6
                                  2251.2
## lme2 8 2267.1 2295.2 -1125.5
                                 2251.1 0.1274
                                                            0.9383
# Output from the random intercept model
summary(lme1)
## Linear mixed model fit by REML ['lmerMod']
## Formula: LMA ~ species + dfromtop + species:dfromtop + (1 | ID)
##
      Data: pref
##
## REML criterion at convergence: 2240
## Scaled residuals:
      Min 1Q Median
                              3Q
                                      Max
## -2.3476 -0.5146 -0.0901 0.4656 3.3215
##
## Random effects:
## Groups Name
                        Variance Std.Dev.
## ID
       (Intercept) 943.4 30.71
## Residual
                        327.2 18.09
## Number of obs: 249, groups: ID, 35
```

```
##
## Fixed effects:
                                 Estimate Std. Error t value
## (Intercept)
                                177.9437 8.0193 22.189
                               134.9772 11.2094 12.041
## speciesPinus ponderosa
## dfromtop
                                -2.0001 0.3030 -6.602
## speciesPinus ponderosa:dfromtop -0.8197 0.5152 -1.591
## Correlation of Fixed Effects:
              (Intr) spcsPp dfrmtp
## spcsPnspndr -0.715
## dfromtop -0.309 0.221
## spcsPpndrs: 0.182 -0.309 -0.588
# Using Anova from car, we get p-values for the main effects.
library(car)
Anova(lme1)
## Analysis of Deviance Table (Type II Wald chisquare tests)
##
## Response: LMA
##
                    Chisq Df Pr(>Chisq)
## species
                 147.475 1
                                <2e-16 ***
                  86.832 1
                                 <2e-16 ***
## dfromtop
## species:dfromtop 2.531 1
                                 0.1116
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

We now conclude that LMA decreases significantly with dfromtop. Compare this with the fixed-effects model we started with:

```
summary(lm1)
## Call:
## lm(formula = LMA ~ species + dfromtop + species:dfromtop, data = pref)
##
## Residuals:
             1Q Median
      Min
                        3Q
                                   Max
## -80.047 -21.737 -2.908 17.231 109.207
## Coefficients:
                               Estimate Std. Error t value Pr(>|t|)
                              164.0749 4.5800 35.824 <2e-16 ***
## (Intercept)
## speciesPinus ponderosa
                              125.2444 6.2581 20.013 <2e-16 ***
                               ## dfromtop
## speciesPinus ponderosa:dfromtop 1.1855 0.6923 1.712 0.0881.
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 31.34 on 245 degrees of freedom
## Multiple R-squared: 0.8211, Adjusted R-squared: 0.8189
## F-statistic: 374.9 on 3 and 245 DF, p-value: < 2.2e-16
```

Ignoring the tree-to-tree variance thus resulted in drawing the wrong conclusion from our data. When we accounted for this variation with a mixed-effects model, we did find a significant overall relationship between

LMA and dfromtop. The reason for this discrepancy is that the large variation in intercepts between the individual trees masked the relationship between the two variables within individuals.

Finally, we may be interested in quantifying the variation between individuals in terms of the intercept and slope. These are the standard deviations of the random effects, and can be extracted with the VarCorr function (it is also shown in the summary statement of the mixed-effects model).

Try this yourself Try using ranef on 1me1 and 1me2; the first will show intercepts for each of the random effects (trees), while the second will show both estimated intercepts and slopes for the random effects. Repeat the above example looking at the relationship between narea (leaf nitrogen per unit area), dfromtop, and species. Fitting a random intercept and slope may result in a convergence error, suggesting a poorly fitting model. If this happens, just ignore this step.

1.3 Example: blocked designs

Blocked designs are often used in field experiments to account for known or suspected environmental gradients at the study site. By blocking the experimental design, the effect of the environmental gradient can be separated from the effect of the treatment(s) of interest increasing the ability to detect significant effects. Let's look at the effects of herbicide and profile on soybean litter decomposition as a function of agricultural management (herbicide usage), microenvironment, and time. In this experiment, herbicide treatments were applied at the level of whole plots, with both treatments represented within each of four blocks. Both levels of variety and profile were each represented within each plot, with six replicates of each treatment added to each plot. The data description can be found in Section ??.

The following code prepares the dataset for analysis, and produces Fig. 1.3.

```
# Read data
litter <- read.csv("masslost.csv")

# Make sure the intended random effects (plot and block) are factors
litter$plot <- as.factor(litter$plot)
litter$block <- as.factor(litter$block)

# Represent date as number of days since the start of the experiment
library(lubridate)
litter$date <- as.Date(mdy(litter$date))
litter$date2 <- litter$date - as.Date("2006-05-23")

# Quickly visualize the data to look for treatment effects
library(lattice)
bwplot(masslost ~ factor(date) | profile:herbicide, data=litter)</pre>
```

From inspecting Fig. 1.3, the buried litter appears to be decomposing faster than the surface litter (masslost is higher for buried compared to surface). If there are effects of herbicide (gly vs. conv), they are not immediately clear from the figure.

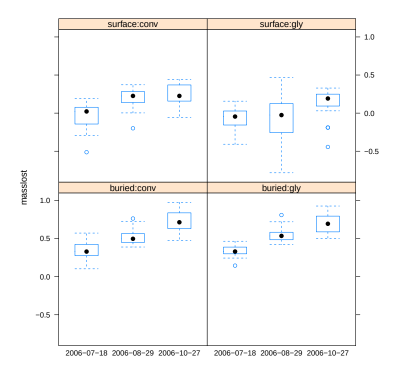


Figure 1.3: Proportion of litter mass lost from bags during field incubation as a function of microenvironment and herbicide program.

The blocking can be treated as a fixed effect or random effect. However, the design is unbalanced because some litter bags were lost, resulting in a variable number of litter bags recovered from each treatment. This affects the calculation of sums of squares, which vary depending on the order the terms are introduced to the model. It is therefore more appropriate to treat the blocking factor as a random effect, and use a mixed-effects model.

We first fit a simple linear model which ignores some details of the experimental design, and use block as a fixed effect. It is often very useful to start with a linear model, perhaps on subsets of the data, to gradually try to make sense of the data.

```
# Count the data to confirm that the design is unbalanced (ignore blocks for brevity)
ftable(xtabs(~ date2 + profile + herbicide, data=litter))
##
                  herbicide conv gly
## date2 profile
## 56
         buried
                              22
                                   22
##
         surface
                              21
                                  21
                              23
                                  23
## 98
         buried
##
         surface
                              20
                                   20
## 157
                              19
                                  16
         buried
##
         surface
                              18
                                  21
# Simple linear model with 'herbicide' as the first predictor in the model,
m1fix <- lm(masslost ~ date2 + herbicide * profile + block, data = litter)</pre>
anova(m1fix)
## Analysis of Variance Table
##
```

```
## Response: masslost
##
                  Df Sum Sq Mean Sq F value
                                                 Pr(>F)
## date2
                    1 3.1696 3.1696 122.0249 < 2.2e-16 ***
## herbicide
                    1 0.4327 0.4327 16.6579 6.113e-05 ***
## profile
                     1 13.7225 13.7225 528.3018 < 2.2e-16 ***
                     3 0.4674 0.1558 5.9987 0.0005891 ***
## block
## herbicide:profile 1 0.3207 0.3207 12.3475 0.0005284 ***
## Residuals 238 6.1820 0.0260
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# ... or listing 'profile' first in the model.
m2fix <- lm(masslost ~ date2 + profile * herbicide + block, data = litter)
anova(m2fix)
## Analysis of Variance Table
## Response: masslost
                    Df Sum Sq Mean Sq F value
## date2
                    1 3.1696 3.1696 122.0249 < 2.2e-16 ***
## profile
                     1 13.8335 13.8335 532.5749 < 2.2e-16 ***
## herbicide
                    1 0.3217 0.3217 12.3848 0.0005184 ***
                    3 0.4674 0.1558 5.9987 0.0005891 ***
## profile:herbicide 1 0.3207 0.3207 12.3475 0.0005284 ***
## Residuals 238 6.1820 0.0260
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

The sums of squares and p-values differ for profile and herbicide across the two fits (although they are still highly significant). Note that the order of the variables entered in the model matters because each next term is tested against a model that includes *all terms preceding it* (so-called Type-I tests). These standard tests with anova are sequential tests, which is perhaps not the most intuitive behaviour.

In many cases it is more intuitive to use so-called Type-II tests, in which each main effect is tested against a model that includes *all other terms*. We can use Anova (from the car package) to do this.

```
library(car)
Anova(m1fix)
## Anova Table (Type II tests)
##
## Response: masslost
                    Sum Sq Df F value
##
                                          Pr(>F)
## date2
                    3.7056 1 142.6601 < 2.2e-16 ***
## herbicide
                   0.3330 1 12.8198 0.0004157 ***
## profile
                  13.6594 1 525.8732 < 2.2e-16 ***
## block
                    0.4933
                            3 6.3311 0.0003793 ***
## herbicide:profile 0.3207 1 12.3475 0.0005284 ***
## Residuals
                  6.1820 238
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Anova(m2fix)
## Anova Table (Type II tests)
## Response: masslost
```

If the data were in fact balanced, the sequential and Type-II tests would be identical.

We also have not yet accounted for the fact that multiple litter bags were placed within single plots and that the 'herbicide' treatment was applied at the level of the plots, not the individual bags, which further complicates the analysis. Treating block and plot as random effects addresses both the imbalance and the hierarchical nature of the design.

In this example, we specify the nested nature of the data (plots within blocks) in the formula for the random effects as (1|block/plot).

```
# fit model with random effects, plots nested within blocks
m1 <- lmer(masslost ~ date2 + herbicide * profile + (1|block/plot),
          data = litter)
Anova(m1)
## Analysis of Deviance Table (Type II Wald chisquare tests)
## Response: masslost
                     Chisq Df Pr(>Chisq)
##
## date2
                  142.525 1 < 2.2e-16 ***
## herbicide
                   12.749 1 0.0003561 ***
## profile 526.246 1 < 2.2e-16 ***
## herbicide:profile 12.187 1 0.0004813 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# As you can see for yourself with anova(m1, m2), plot explains
# essentially zero variance.
m2 <- lmer(masslost ~ date2 + herbicide * profile + (1|block),</pre>
          data = litter)
Anova(m2)
## Analysis of Deviance Table (Type II Wald chisquare tests)
## Response: masslost
                     Chisq Df Pr(>Chisq)
##
                  142.525 1 < 2.2e-16 ***
## date2
## herbicide
## profile
                   12.749 1 0.0003561 ***
                    526.246 1 < 2.2e-16 ***
## herbicide:profile 12.187 1 0.0004813 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Note that in the model, we used date2 as a numeric variable, which assumes that the relationship between masslost and date2 is more or less linear. Figure 1.4 shows that this is a resonable assumption However, in the case where you have a timeseries where no transformation exists to linearize the relationship, you will have to represent your time variable as a factor. We return to this issue in Section 1.5, after we treat another

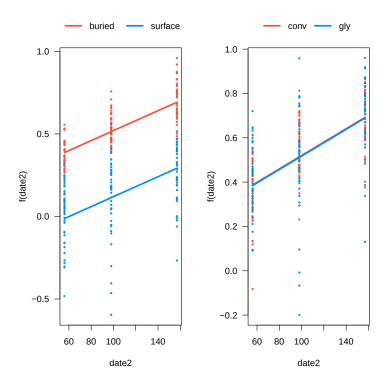


Figure 1.4: Visualization of the fixed effects of the mixed-effects model fit to the litter decomposition data.

repeated measures example where time was continuous in Section 1.5.

Finally we visualize the fit, to make sense of the fitted coefficients, and to make sure we draw the right conclusions as to the direction of the significant effects. As we saw in Chapter ??, we can use visreg to quickly visualize the fitted model.

```
# Because we have three fixed effects, we can make two plots to visualize all combinations.
par(mfrow=c(1,2))
visreg(m2, "date2", by="profile", overlay=TRUE)
visreg(m2, "date2", by="herbicide", overlay=TRUE)
```

1.3.1 More about p-values

As we mentioned at the top of this chapter, p-values in lme4 are a bit controversial. We here used the Anova function, but another option is to use the LMERConvenienceFunctions package. This package includes functions to estimate of p-values from *conservative* and *liberal* assumptions about random effect degrees of freedom, and thus gives a range of possible p-values.

```
library(LMERConvenienceFunctions)

# To use the function below, we must fit with ML, not REML.
m2ml <- update(m2, REML=FALSE)

# calculate upper- and lower-bounds on p-values
pamer.fnc(m2ml)</pre>
```

```
##
                   Df Sum Sq Mean Sq F value upper.den.df upper.p.val
## date2
                    1 3.1860 3.1860 124.7135
## herbicide
                    1 0.4439 0.4439 17.3742
                                                      241
                                                                0e+00
## profile
                    1 13.6726 13.6726 535.2013
                                                      241
                                                                0e+00
## herbicide:profile 1 0.3153 0.3153 12.3406
                                                                5e-04
                   lower.den.df lower.p.val expl.dev.(%)
## date2
                            237
                                     0e+00
## herbicide
                            237
                                     0e+00
                                                1.8269
## profile
                            237
                                     0e+00
                                                56.2776
## herbicide:profile
                                     5e-04
                                                1.2976
                            237
```

Each main effect and interaction has two p-values: one assuming that each random effect accounts for one degree of freedom (lower.p.val) or no degrees of freedom (upper.p.value). The 'true' p-value will be somewhere in between these two bounds.

In this particular case, the results are highly significant (both lower and upper p-values are very small) because the effect size is quite large, but this will not always be the case.

Try this yourself Use pamer.fnc on the model above that contains random effects for both Block and Plot. What effect does this have on the degrees of freedom and p-value calculations? Why?

Another approach is to evaluate the importance of a term by comparing models that contain or do not contain that term using likelihood ratio tests with anova (p-values are approximate since likelihood ratios don't quite fit a chi-square distribution) or model selection based on AIC (the lower the better).

```
# remove the interaction term from the model
m2.int <- lmer(masslost ~ date2 + herbicide + profile + (1|block), data = litter)
# compare the two models
# Note that anova() will refit the models with ML (not REML) automatically,
# this is necessary when comparing models with different fixed or random effects terms.
anova(m2m1, m2.int)
## refitting model(s) with ML (instead of REML)
## Data: litter
## Models:
## m2.int: masslost ~ date2 + herbicide + profile + (1 | block)
## m2ml: masslost ~ date2 + herbicide * profile + (1 | block)
               AIC
                        BIC logLik deviance Chisq Chi Df Pr(>Chisq)
## m2.int 6 -173.68 -152.64 92.838 -185.68
        7 -183.70 -159.16 98.848 -197.70 12.02
## m2m1
                                                     1 0.0005263 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
AIC(m2m1, m2.int)
##
         df
                  AIC
          7 -183.6958
## m2m1
## m2.int 6 -141.5371
```

The model that includes the interaction provides the much better model fit. We can tell this by the significant p-value from the anova result and by the lower AIC score for the model that includes an interaction. When an interaction is significant, the automatic follow-up question is 'what is the source of this interaction?'. Again inspecting Fig. 1.4, it appears bthat the herbicide treatments affected decomposition differently, but only on the surface of the soil. To further understand the nature of the interactions, it is useful to combine variables into a single variable, as the following example illustrates.

```
# Create a new variable containing the combinations of herbicide and profile.
# This new variable will have 4 levels
litter$combtrt <- paste(litter$herbicide, litter$profile, sep='-')</pre>
# Lump all observations for which the bags were buried into a single level,
# we now have just three levels in the new combined variable.
litter$combtrt[litter$profile == 'buried'] <- 'buried'</pre>
# Make this new variable into a factor
litter$combtrt <- as.factor(litter$combtrt)</pre>
# Fit a models using this new factor (3 levels) and a model without herbicide (2 levels)
m3 <- lmer(masslost ~ date2 + combtrt + (1|block), data = litter, REML=FALSE)
m3.herb <- lmer(masslost ~ date2 + profile + (1|block), data = litter, REML=FALSE)
# Compare the models by AIC (lower is 'better')
AIC(m2ml, m3, m3.herb)
         df
           7 -183.6958
## m2m1
            6 -185,6860
## m3.herb 5 -163.6908
```

The model with the lowest AIC is m3, which is the model describing the relationship where herbicide affected decomposition rates only at the soil surface (because for that model, we combined all 'buried' litter samples into one level, regardless of the herbicide application).

1.4 Example: repeated measures

This example shows a very common use of mixed-effects models in repeated measurements. The basic idea is that when you have measurements on the same individuals (or plots, or some other unit) over time, you cannot treat the measurements as independent because that would be pseudo-replication, inflation of your sample size, and anti-conservative conclusions about significant effects.

We use data from the Hawkesbury Forest Experiment irrigation by fertilisation experiment (HFEIF, see Section ?? for description of the data). In this experiment, sixteen plots of 72 *Eucalyptus saligna* trees were remeasured 20 times for height and diameter (although on a number of dates, not all trees were measured). Four treatments were applied (control, irrigated, fertilised, irrigated + fertilised). We ask in the following example whether tree height differs by treatment.

It is important that you recognize that the experimental unit in this example is the plot, not the tree, because the treatments were applied at a plot level. We therefore have to take into account the fact that trees are nested in plots, to avoid pseudoreplication.

```
# Read data, make proper date and make sure the intended factor variables are factors.
hfeif <- read.csv("HFEIFbytree.csv")
hfeif$Date <- as.Date(hfeif$Date)

# Make sure plot number (plotnr) is a factor; it is read in as a numeric variable.
hfeif$plotnr <- as.factor(hfeif$plotnr)

# Days since start of experiment
# The as.numeric statement converts this into a simple numeric variable
hfeif$Time <- as.numeric(with(hfeif, Date - min(Date)))</pre>
```

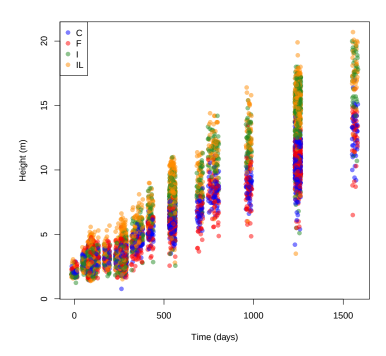


Figure 1.5: Plot tree height over time for the HFE irrigation x fertilisation experiment.

Before we do anything, always explore the data with a few simple figures. Here we show tree height over time, colored by treatment (Fig. 1.5). The data show some separation between at least some of the treatments over time. Also note that the increase in height over time is perhaps not exactly linear, but we will ignore this in the remainder of the example (and further note that no straightforward transformation exists in this case).

Note the use of <code>jitter</code> in the example below, this adds some random noise to the Time variable to avoid excessive overlap of data points on each Date. We also use <code>sample</code> to randomly reorder the rows of the dataset to avoid the final treatment in the dataset to be plotted on top (this way, it is easier to see treatment differences).

The following code produces Fig. 1.5.

Again, the reason we want to use mixed-effects models in this case is because we want to use the correct number of degrees of freedom to test for the treatment effect. If you are not sure what that should be, let's start with a linear model on the data from just one date, when we have averaged the data by plot (our experimental unit). In this case we can simply use lm, as follows.

The following example produces Fig. 1.6.

```
# Take subset of data at last Date
hfeif_last <- subset(hfeif, Date == max(Date))</pre>
# Average all variables by plot (and include the 'treat' factor variable in the result)
library(doBy)
hfeif_last_plot <- summaryBy(. ~ Date + plotnr, data=hfeif_last,</pre>
                             FUN=mean. na.rm=TRUE.
                             id=~treat, keep.names=TRUE)
# Linear model with treatment only.
lm_last <- lm(height ~ treat, data=hfeif_last_plot)</pre>
# Note that height is highly significant, and that we use 3 numerator df
# to test for treatment effects
anova(lm_last)
## Analysis of Variance Table
##
## Response: height
            Df Sum Sq Mean Sq F value
            3 74.160 24.7201 19.137 7.226e-05 ***
## treat
## Residuals 12 15.501 1.2917
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# A quick visualization of the fitted model shows much taller trees
# in I and IL,
library(visreg)
visreg(lm_last, "treat", xlab="Treatment", ylab="Tree height (m)")
```

The above shows 3 numerator degrees of freedom in the F-test, which makes sense because we have 4 levels of our treatment (thus df = 4 - 1). You can do no such simple check for the denominator degrees of freedom, but it's a useful check nonetheless.

To account for the repeated measures nature of the data as well as the fact that the experimental unit is the plot, not the tree, all we need is to specify the plot as the random effect. We will fit two models, one without and one with the interaction between Time and treat, and again use Anova (from the car package) to test for significant effects.

Note that we specify Time as a numeric variable, which assumes that the relationship between height and Time is more or less linear, which according to Figure 1.5 is a reasonable assumption. We return to this issue in Section 1.5.

```
# Effect of treatment on intercept only.
lmeif1 <- lmer(height ~ treat + Time + (1|plotnr), data=hfeif)
Anova(lmeif1)

## Analysis of Deviance Table (Type II Wald chisquare tests)

##
## Response: height

## Chisq Df Pr(>Chisq)

## treat 27.752 3 4.095e-06 ***

## Time 48339.942 1 < 2.2e-16 ***

## ---

## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

# Effect of treatment on intercept and slope (i.e. main effect + interaction)</pre>
```

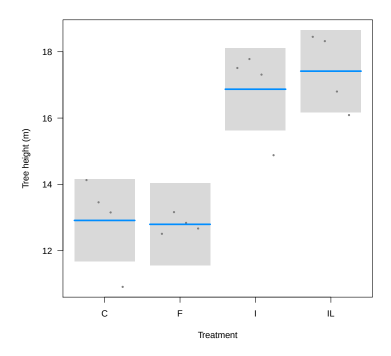


Figure 1.6: Simple visualization of fitted linear model (with lm) of tree height on the last date of the HFE IF data.

```
lmeif2 <- lmer(height ~ treat*Time + (1|plotnr), data=hfeif)</pre>
Anova(lmeif2)
## Analysis of Deviance Table (Type II Wald chisquare tests)
##
## Response: height
##
                 Chisq Df Pr(>Chisq)
                27.389 3 4.879e-06 ***
## treat
             72342.647 1 < 2.2e-16 ***
## Time
## treat:Time 3208.442 3 < 2.2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# A likelihood ratio tests shows the interaction is highly significant
anova(lmeif1, lmeif2)
## refitting model(s) with ML (instead of REML)
## Data: hfeif
## Models:
## lmeif1: height ~ treat + Time + (1 | plotnr)
## lmeif2: height ~ treat * Time + (1 | plotnr)
              AIC
                   BIC logLik deviance Chisq Chi Df Pr(>Chisq)
## lmeif1 7 22654 22702 -11320
                                  22640
## lmeif2 10 20054 20122 -10017
                                  20034 2605.9
                                                    3 < 2.2e-16 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

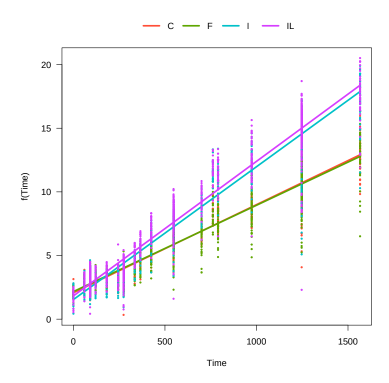


Figure 1.7: Visualized effect of Time on height, by treatment for the HFE IxF dataset, fitted with a linear mixed-effects model.

Note that in the above, we have correctly used 3 numerator degrees of freedom to test for the effect of treatment on height. The individual Anova statements summarize the significance of the fixed effects in each model, whereas the anova of the two models uses a likelihood-ratio test on the two models. In this case, it effectively tests for significance of the interaction (because the only difference between the two models was the inclusion of the treat by Time interaction in lmeif2). The interaction is overwhelmingly significant.

The final step is to try to understand this interaction, how large is the effect size, and which direction does it point? The time by treatment interaction is significant, but in which way? It is never sufficient in an analysis to state that an interaction was 'significant', we must make more sense of it. One simple approach is to use the visreg package to visualize the fit (see Fig. 1.7).

```
library(visreg)
visreg(lmeif2, "Time", by="treat", overlay=TRUE)
```

In this case it is abundantly clear that irrigated (I) and irrigated + fertilised (IL) have a steeper slope of height with Time (that is, they have a faster height growth), compared to control (C) and fertilised (F). If there was no significant interaction (or a small effect size), the lines would be parallel to each other.

We can further look at the p-values for the individual effects (slopes and intercepts by treatment). Note that p-values in the summary statement are only computed if we have loaded the lmerTest package before fitting the model. Consider this example,

```
# Loading this package first affects both summary and anova methods
library(lmerTest)
##
## Attaching package: 'lmerTest'
```

```
## The following object is masked from 'package:lme4':
##
##
      lmer
## The following object is masked from 'package:stats':
##
##
      step
# ... we must refit the model after loading lmerTest
lmeif2 <- lmer(height ~ treat*Time + (1|plotnr), data=hfeif)</pre>
# Print just the coefficients table from the summary
summary(lmeif2)$coefficients
##
                                                                    Pr(>|t|)
                   Estimate
                              Std. Error
                                                 df
                                                        t value
## (Intercept) 2.073764669 2.992535e-01 12.32252
                                                      6.9297919 1.377196e-05
               0.093074081 4.233121e-01 12.33460 0.2198711 8.295684e-01
## treatF
               -0.535359455 4.232604e-01 12.32858 -1.2648466 2.293128e-01
## treatI
              -0.246322580 4.232390e-01 12.32609 -0.5819940 5.710624e-01
## treatIL
               0.006921244 6.399533e-05 6453.01255 108.1523211 0.000000e+00
## treatF:Time -0.000141759 9.111223e-05 6453.02451 -1.5558723 1.197875e-01
## treatI:Time 0.003514427 9.095049e-05 6453.01005 38.6411001 0.000000e+00
## treatIL:Time 0.003645960 9.085136e-05 6453.02770 40.1310445 0.000000e+00
```

Try this yourself The lmerTest package also modifies the anova function, so that it calculates p-values for a fitted model with lmer. Compare anova(lmeif2) with Anova(lmeif2), these will rarely be exactly the same as they use different methods to approximate the degrees of freedom of the random effects.

Looking at the interaction terms (treat:Time), the summary table shows that treatF:Time is not significantly different from the first level (treatC:Time), that is, there is no difference between fertilized and control in terms of the interaction with Time. But both irrigated (I) and irrigated + fertilised (IF) are highly significant, again, this comparison is in relation to the first level of the factor (control, C).

Although there is a significant main effect of treatment, none of the levels are actually different from the first (the control). This shows that the intercept itself is different from zero, but the treatments are not actually different in terms of the intercept. This makes sense, because seedlings were planted at time zero before any treatment was applied.

1.5 Repeated measures: is time numeric or factor?

In both examples in this chapter where we used time as a predictor in our models, we treated time as a continuous (numeric) variable. This was appropriate in both cases because the relationship between the dependent variable (masslost or height) showed a nearly linear relationship with time, allowing us to estimate and interpret an intercept and a slope of the variable with time. In the example with the tree height measurements, the slope of height with time can actually be interpreted as the height growth rate.

But there are many cases in which it would be more appropriate to use time as a factor variable. These include cases where the relationship is highly non-linear and cannot be transformed, or you only have two or three dates of measurements. The example below shows a simple example for the latter case. (see also Fig. 1.8).

```
# A repeated measures example with only two dates of measurement.
# Though it is possible to have time as a continuous variable, it is much
```

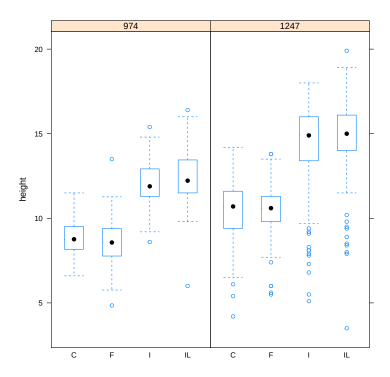


Figure 1.8: Boxplots of height vs. treatment and time (days since start of experiment, shown in the panel label) for a subset of the HFE IF data.

```
# more useful to code it as a factor.

# We take a subset of hfeif.
hfeif2 <- subset(hfeif, Date %in% as.Date(c("2010-09-01","2011-06-01")))

# Convert Time to a factor
hfeif2$Time_fac <- as.factor(hfeif2$Time)

# As before, we can quickly use bwplot to inspect the data
library(lattice)
bwplot(height ~ treat | Time_fac, data=hfeif2)</pre>
```

Try this yourself Repeat the above example, but using the entire dataset (rather than a subset of the data for two dates). Inspect the model with Anova, and also look at the summary statement.

Let's fit the mixed-effects model on this small subset of the data to test whether a) treatment affects height, b) there is an effect of time on height, c) there is an interaction (i.e. height response to treatment depends on time).

```
# Fit the model
lmeif4 <- lmer(height ~ treat*Time_fac + (1|plotnr), data=hfeif2)

# Overall significance shows no interaction
Anova(lmeif4)

## Analysis of Deviance Table (Type II Wald chisquare tests)</pre>
```

```
##
## Response: height
## Chisq Df Pr(>Chisq)
## treat 69.0334 3 6.874e-15 ***
## Time_fac 359.1303 1 < 2.2e-16 ***
## treat:Time_fac 8.4144 3 0.03818 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1</pre>
```

Try this yourself Use visreg to visualize the fit, and compare it to the box plots produced above.

1.6 Generalized linear mixed effects models

We have seen how to fit generalised linear models (section ??) and linear mixed models (above). Once you know how to fit these models in **R**, fitting GLMMs is fairly easy using the glmer function in the lme4 package.

The EucFACE ground cover dataset (see Section $\ref{section}$) contains estimates of plant and litter cover within the rings of the EucFACE experiment, evaluating forest ecosystem responses to elevated CO_2 , on two dates. There are six rings (Ring), three for each treatment (Trt; ambient and elevated CO_2). Within each ring are four plots (Plot) and within each plot are four 1m by 1m subplots (Sub). Here we will test for an interaction between Trt and Date on ground cover measurements of Forbes (these are count data).

The following code produces Fig. 1.9, showing a plot of the raw groundcover data.

```
# read data and convert random effects to factors
eucface <- read.csv("eucfaceGC.csv")
eucface$Ring <- as.factor(paste(eucface$Ring, eucface$Trt, sep='-'))
eucface$Plot <- as.factor(eucface$Plot)
eucface$Sub <- as.factor(eucface$Sub)

# load packages
library(lme4)
library(lattice)

# A quick plot to visualize ground cover by Date and Ring.
# Colours represent 'Plot'.
xyplot(Forbes-Date|Ring,groups=Plot,data=eucface,pch=16,jitter.x=T)</pre>
```

Since the data are count data, it is usually appropriate to use the Poisson distribution. The following code fits a glmer with the poisson error family, and produces diagnostic plots are in Fig. ??.

To usual way to decide on the appropriate family is to inspect the diagnostic plots, especially a plot of residuals versus the fitted values. In the following, we fit the model three times, first with normal errors (using lmer), then with Poisson errors (the distribution we expect to be appropriate), and a Poisson distribution with square-root link function.

The following code produces three diagnostic plots (Fig. 1.10).

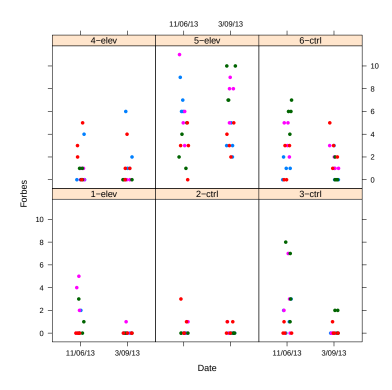


Figure 1.9: Counts of forbs within plant communities exposed to ambient or elevated carbon dioxide concentrations at two dates. Points represent estimates within subplots, subplots common to the same plot within each ring have a common colour.

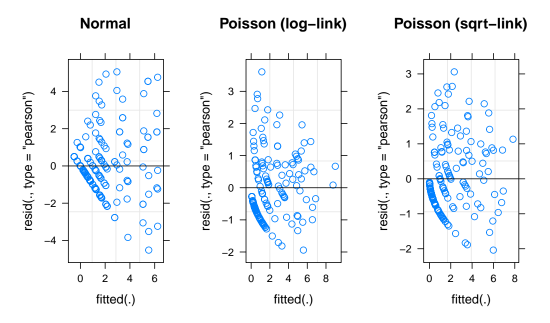


Figure 1.10: Diagnostic plots for three model fits of the EucFACE groundcover data.

Note that the syntax of glmer is identical to that of lmer, with the exception of the family argument.

Inspecting Fig. 1.10, the residuals look better for the models fit with the Poisson family compared to the gaussian error (as assumed by lmer). The residuals improve a bit more when using the sqrt link function in the Poisson family.

Note that the fit of the model may be improved further by using one of the other plant cover variables as a covariate or by incorporating other data from the site, but for the purpose of this chapter the fit is good enough.

Something else to consider when it comes to poisson errors is that overdispersion can result in the underestimation of error terms for the model coefficients. Overdispersion occurs when you have a large number of zeros in the data and/or an important predictor is not accounted for. Testing hypotheses from models where overdispersion is evident is dangerous as the probability of Type I error is increased. A good model fit should result in the ratio of residual deviance to degrees of freedom being close to one.

```
# Calculate the residual deviance
sum(resid(forb.pois.sqrt, type='pearson')^2)
## [1] 171.408
# model summary, note 'df.resid' in the model summary is 185
summary(forb.pois.sqrt)
```

```
## Generalized linear mixed model fit by maximum likelihood (Laplace
    Approximation) [glmerMod]
##
  Family: poisson (sqrt)
## Formula: Forbes ~ Date * Trt + (1 | Ring/Plot/Sub)
##
     Data: eucface
##
##
       ATC
               BIC
                      logLik deviance df.resid
##
     612.6
              635.4
                      -299.3
                               598.6
                                          185
##
## Scaled residuals:
##
      Min
              1Q Median
                              ЗQ
                                     Max
## -2.0391 -0.5597 -0.2764 0.4883 3.0589
##
## Random effects:
## Groups
                              Variance Std.Dev.
                   Name
## Sub:(Plot:Ring) (Intercept) 0.05631 0.2373
## Plot:Ring
                   (Intercept) 0.12549 0.3542
## Ring
                   (Intercept) 0.32556 0.5706
## Number of obs: 192, groups: Sub:(Plot:Ring), 96; Plot:Ring, 24; Ring, 6
##
## Fixed effects:
##
                      Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                       ## Date3/09/13
                       -0.5541
                                  0.1132 -4.894 9.88e-07 ***
## Trtelev
                       0.1218
                                  0.5022 0.243 0.80833
                                  0.1604 2.325 0.02007 *
## Date3/09/13:Trtelev
                      0.3729
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
              (Intr) Dt3/09/13 Trtelv
## Date3/09/13 -0.152
             -0.707 0.109
## Trtelev
## Dt3/09/13:T 0.106 -0.710
                              -0.150
```

Overdispersion does not appear to be a problem here as the ratio is less than one. If it was, we could use a quasipoisson family as for GLMs (but unfortunately glmer does not support that family). It has also been suggested that including individual-level random effects in the model could alleviate the problem of overdispersion. We try this in the following example.

```
# Create a variable for individual-level random effects
eucface$Ind <- as.factor(1:nrow(eucface))</pre>
# fit the model and look at the model summary
forb.pois.ind <- glmer(Forbes~Date*Trt+(1|Ring/Plot/Sub/Ind),</pre>
                        family=poisson(sqrt), data=eucface)
summary(forb.pois.ind)
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
##
## Family: poisson ( sqrt )
## Formula: Forbes ~ Date * Trt + (1 | Ring/Plot/Sub/Ind)
##
      Data: eucface
##
##
        AIC
                 BIC
                        logLik deviance df.resid
```

```
##
       612
               638
                      -298
                               596
                                        184
##
## Scaled residuals:
      Min 1Q Median
##
                            ЗQ
                                   Max
## -1.7030 -0.5273 -0.2436 0.4350 2.5908
## Random effects:
  Groups
                       Name
                                  Variance Std.Dev.
##
## Ind:(Sub:(Plot:Ring)) (Intercept) 0.06872 0.2621
## Sub:(Plot:Ring) (Intercept) 0.01199 0.1095
## Plot:Ring
                       (Intercept) 0.12407 0.3522
                       (Intercept) 0.32530 0.5703
## Ring
## Number of obs: 192, groups:
## Ind:(Sub:(Plot:Ring)), 192; Sub:(Plot:Ring), 96; Plot:Ring, 24; Ring, 6
##
## Fixed effects:
                    Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                     1.1766 0.3564 3.302 0.000961 ***
## Date3/09/13
                    0.1219
## Trtelev
                                0.5031 0.242 0.808482
## Date3/09/13:Trtelev 0.3763
                                0.1788 2.104 0.035368 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
##
             (Intr) Dt3/09/13 Trtelv
## Date3/09/13 -0.170
## Trtelev -0.706 0.121
## Dt3/09/13:T 0.118 -0.710
                            -0.168
```

The random effects block indicates that a small amount of variance is accounted for by the individual level random effects. This is expected as we did not observe overdispersion.

We can use Anova function from the car package to calculate significance associated with the main effects and interaction. Note that is is only one way of many to calculate p-values, as we discussed above linear mixed models.

There appears to be a significant interaction between Date and Trt. Again, when an interaction is significant we must dig deeper to understand the source of the interaction. Looking at Fig. 1.11, it appears that:

- 1. Forbs decreased in frequency between the two dates in the control but not in the elevated ${\rm CO_2}$ treatment
- 2. Forb abundance was lower in the control than in the elevated CO₂ treatment on the second date, but

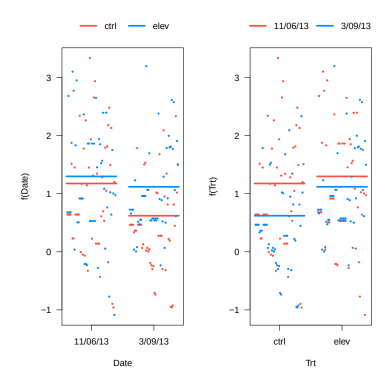


Figure 1.11: Model predictions, looking at treatment effects by date (left) and date effects by treatment (right).

not the first date.

```
# set up graphics window for two plots
par(mfrow=c(1, 2))

# plot model preditions and data
visreg(forb.pois.ind, 'Date', 'Trt', overlay=TRUE)
visreg(forb.pois.ind, 'Trt', 'Date', overlay=TRUE)
```

We can evaluate each of these hypotheses through model selection after combining treatment levels.

```
# create a three level factor that combines both dates in the 'elev' treatment
eucface$trtcomb.elev <- with(eucface, paste(Trt, Date, sep='-'))</pre>
eucface$trtcomb.elev[eucface$Trt == 'elev'] <- 'elev'</pre>
eucface$trtcomb.elev <- as.factor(eucface$trtcomb.elev)</pre>
levels(eucface$trtcomb.elev)
## [1] "ctrl-11/06/13" "ctrl-3/09/13" "elev"
# create a three level factor that combines both treatments for the '11/06/13' sampling
eucface$trtcomb.date <- with(eucface, paste(Trt, Date, sep='-'))</pre>
eucfacetrtcomb.date[eucface\\Date == '11/06/13'] <- '11/06/13'
eucface$trtcomb.date <- as.factor(eucface$trtcomb.date)</pre>
levels(eucface$trtcomb.date)
## [1] "11/06/13"
                       "ctrl-3/09/13" "elev-3/09/13"
# fit model with three level combination factors
m2.elev <- glmer(Forbes ~ trtcomb.elev + (1|Ring/Plot/Sub/Ind),</pre>
                  family=poisson('sqrt'), data=eucface)
```

```
m2.date <- glmer(Forbes ~ trtcomb.date + (1|Ring/Plot/Sub/Ind),
                 family=poisson('sqrt'), data=eucface)
# fit models with only one main effect (Date or Trt)
m3.Trt <- glmer(Forbes~Date+(1|Ring/Plot/Sub/Ind), family=poisson(sqrt), data=eucface)
m3.Date <- glmer(Forbes~Trt+(1|Ring/Plot/Sub/Ind), family=poisson(sqrt), data=eucface)
# compare models (model with the lowest AIC is the most efficient at predicting the response)
AIC(forb.pois.ind, m2.elev, m2.date, m3.Trt, m3.Date)
                 df
## forb.pois.ind 8 611.9524
## m2.elev
                 7 611.9420
## m2.date
                 7 610.0109
                 6 612.6342
## m3.Trt
## m3.Date
                 6 627,4332
```

The three-level model in which date effects are estimated in the control treatment but not the elevated treatment (m2.elev) is not a substantial improvement over the fully factorial model (forb.pois.ind), so we do not consider it further. The three-level model in which treatment effects are estimated on the second date but not the first date (m2.date) has the lowest AIC score in comparison to the fully factorial model (forb.pois.ind) and to the two-level models that do not estimate an effect of treatment (m3.Trt) or date (m3.Date), so is the model that we select as the one that best predicts forb dynamics.

1.6.1 Logistic regression using mixed effects models

Something that wasn't mentioned regarding this EucFACE ground cover dataset is that vegetation was assessed at a maximum of 16 points within each subplot and, therefore, the maximum number of observations per plot is constrained. This does not affect our analysis of forb abundances because these are generally low (less than ten in almost all plots) and so interpreting these as count data is appropriate. This is not the case for other response variables. As an example, let's look at grass cover in Fig. 1.12.

```
# A quick plot to visualize ground cover by Date and Ring.
# Colours represent 'Plot'.
xyplot(Grass~Date|Ring,groups=Plot,data=eucface,pch=16,jitter.x=T)
```

The data are clearly bounded at both the lower and upper range. We can treat these as binomial distributed, with the presence or absence of grass assessed at each of the sixteen points within each subplot. Do do this, we use cbind to create a two-column response matrix indicating the number of presences and absences within each subplot, as we did for logistic regression in section ??. We also specify the random effects as we did for the analysis of forb abundance above.

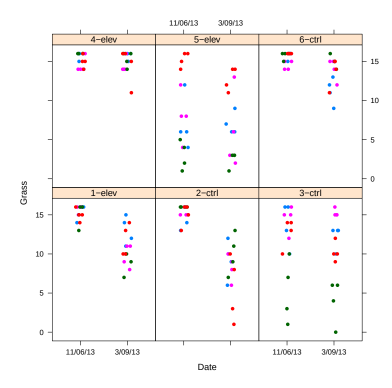


Figure 1.12: Grass cover within plant communities exposed to ambient or elevated carbon dioxide concentrations at two dates. Cover was assessed based on presence at each of sixteen locations within a subplot. Points represent estimates within subplots, subplots common to the same plot within each ring have a common colour.

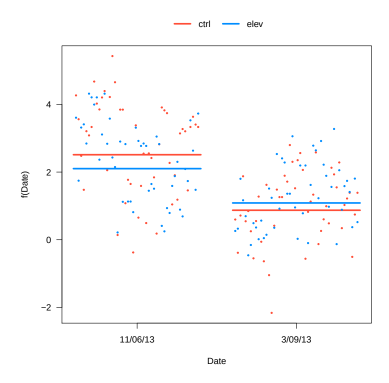


Figure 1.13: Model predictions, looking at treatment effects by date.

```
## Trt     0.0000     1     0.997748
## Date:Trt     8.3183     1     0.003925 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

The results suggest a highly significant effect of date and a significant date by treatment interaction. The model predictions are shown in Fig. 1.13. A similar approach could be used to tease apart the source of the interaction as was used for forb abundances.

```
# plot model preditions and data
visreg(grass.binom, 'Date', 'Trt', overlay=TRUE)
```

1.7 Exercises

In the exercises, we use the following colour codes:

- **Easy**: make sure you complete some of these before moving on. These exercises will follow examples in the text very closely.
- ♦ Intermediate: a bit harder. You will often have to combine functions to solve the exercise in two steps.
- ▲ **Hard**: difficult exercises! These exercises will require multiple steps, and significant departure from examples in the text.

1.7.1 PREF Canopy data

1. ♦ In the analysis of the pref data, use model selection (AIC, anova) to evaluate the importance of species and dfromtop.

1.7.2 Litter decomposition data

1. The litter data contain a factor (variety) describing whether the litter is derived from a genetically modified (gm) or conventional (nongm) soy variety. Plot the data to observe the effect of variety. Use lmer to test the effect of variety, in addition to the other significant variables, on litter decomposition.

1.7.3 EucFACE ground cover data

The file eucfaceGC.csv contains estimates of plant and litter cover within the rings of the EucFACE experiment, evaluating forest ecosystem responses to elevated CO2, on two dates; the data description can be found in Section ?? (p. ??).

- 1. Convert the variables indicating the nested sampling design to factors, then use glmer in lme4 to test for an interaction between Trt and Date on Grass and Litter cover. Grass cover represents a frequency across a maximum of 16 points within a quadrat (use the binomial family), while litter cover represents counts (use the poisson family).
- 2. A Following on from exercise 3, generate subsets to determine the sources of the interactions (i.e., does the treatment effect differ between the two dates or does the date effect differ between the two treatments?).

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