

## Mixed Models

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## The Big Picture

- Mixed models contain a combination of fixed and random effects.
- It is important to properly identify which variables should be modeled as random and which as fixed.
- In the case of completely balanced designs, ANOVA tables can be used for inference with  $F$  tests, but care must be taken to compute p-values properly.
- Blocking variables are a special type of categorical variable.
- Blocks can be random or fixed depending on the context.

## Parameter Estimation

- Recall the Antigua Corn data from last time.
- The model had three parameters, a mean harvest  $\hat{m}u = 4.2917$ , a site variance  $\hat{\sigma}_s^2 = 2.3677$ , and an individual variance  $\hat{\sigma}_e^2 = 0.5775$ .
- The function `mcmcscamp` provides a means to estimate uncertainty in parameter estimates in mixed models by producing a sample of the parameter values from a Bayesian posterior distribution.
- The 0.025 and 0.975 quantiles of the sample are estimates of a 95% *credible region* which you can interpret as a confidence interval.
- The function estimates logarithms of the variance components, so we need to take exponentials to put them back on scale.
- Credible regions need not be symmetric — for variance components they usually are not.

## MCMC Sample Example

```
> library(DAAG)
> library(lme4)
> corn1.lmer = lmer(harvwt ~ 1 + (1 | site), data = ant111b)
> set.seed(3425)
> corn1.samp = mcmcscamp(corn1.lmer, n = 1000)
> apply(corn1.samp, 2, function(x) quantile(x, probs = c(0.025,
+ 0.975)))

      (Intercept) log(sigma^2) log(site.(ln))
2.5%   2.875464  -1.05292374  -0.01928132
97.5%   5.679904   0.06247114   2.37010691

> apply(corn1.samp, 2, function(x) quantile(exp(x), probs = c(0.025,
+ 0.975)))

      (Intercept) log(sigma^2) log(site.(ln))
2.5%   17.73366   0.3489163   0.9809044
97.5%  292.92262   1.0644637  10.6986503
```

- `set.seed` is unnecessary, but makes the sample repeatable.
- The result of `mcmcscamp` is a matrix with 1000 rows and three columns.
- The `apply` function applies to the columns (2nd argument 2) of `corn1.samp` (1st argument) the function (3rd argument) which find quantiles of either  $x$  or  $\exp(x)$  in this case.

## Credible Intervals

- The estimated intercept is  $\hat{\mu} = 4.29$ ,  $P\{2.88 < \mu < 5.68\} = 0.95$ .
- The estimated site variance is  $\hat{\sigma}_a^2 = 2.37$ ,  $P\{0.98 < \sigma_a^2 < 10.7\} = 0.95$ .
- The estimated individual variance is  $\hat{\sigma}_e^2 = 0.58$ ,  $P\{0.35 < \sigma_e^2 < 1.06\} = 0.95$ .

## Soil Moisture

- In an experiment, four different irrigation methods are compared.
- There are 16 different plots in the experiment, considered to be a sample from a larger population of plots.
- Each irrigation treatment is used on four plots, assigned at random.
- Each plot is subsampled — three soil moisture measurements are taken within each plot.
- There is one fixed effect — *irrigation treatment*.
- There are two sources of variation — *plot-to-plot* variability and *within-plot* variability.

## Model

- A model is

$$y_{ijk} = \mu + \alpha_i + b_{ij} + e_{ijk}$$

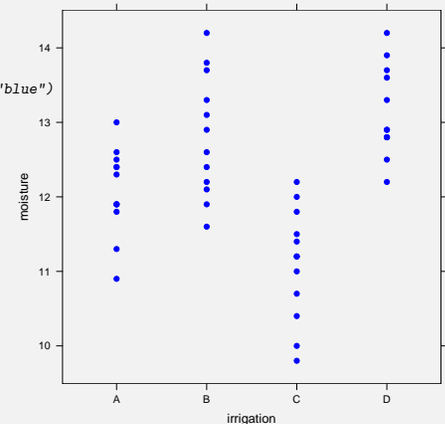
where:

- $\mu$  is an overall mean,
- $\alpha_i$  is the effect of treatment  $i$  where  $\sum_i \alpha_i = 0$ , for  $i = 1, \dots, 4$ ,
- $b_{ij} \sim \text{iid } N(0, \sigma_b^2)$  is the effect of the  $j$ th plot (nested within the  $i$ th treatment),  $j = 1, \dots, 4$ ,
- and  $e_{ijk} \sim \text{iid } N(0, \sigma_e^2)$  is individual random error,  $k = 1, \dots, 3$ .

## Plot of Data

```
> soil = read.table("soil.txt", header = T)
> attach(soil)
> library(lattice)
> fig1 = xyplot(moisture ~ irrigation, pch = 16, col = "blue")
> print(fig1)
```

- Notice that the variation within treatments is substantial, but the treatment distributions appear to be different.
- Variation within treatments looks to be reasonably symmetric without outliers.



## Fitting a Random Effects Model

```

> library(lme4)
> soil.lmer = lmer(moisture ~ irrigation + (1 | irrigation:plot))
> summary(soil.lmer)

```

Linear mixed-effects model fit by REML  
 Formula: moisture ~ irrigation + (1 | irrigation:plot)  
 AIC BIC logLik MLdeviance REMLdeviance  
 83.5 92.86 -36.75 72.22 73.5

Random effects:

Groups	Name	Variance	Std.Dev.
irrigation:plot	(Intercept)	0.44020	0.66347
Residual		0.12813	0.35795

number of obs: 48, groups: irrigation:plot, 16

Fixed effects:

	Estimate	Std. Error	t value
(Intercept)	12.0750	0.3475	34.75
irrigationB	0.7417	0.4914	1.51
irrigationC	-0.9750	0.4914	-1.98
irrigationD	1.0583	0.4914	2.15

Correlation of Fixed Effects:

(Intr)	irrgtB	irrgtC	
irrigationB	-0.707		
irrigationC	-0.707	0.500	
irrigationD	-0.707	0.500	0.500

- irrigation is the fixed effect.
- irrigation:plot is an interaction between irrigation and plot, which is how we specify nesting here.
- There is no main plot effect, so plot does not appear without irrigation in the model.
- The term (1 | irrigation:plot) means that there is a random effect for each plot and this effect is nested within the intercept (the whole model).
- There are two sources of random variation, one for plot and one for individual measurements.

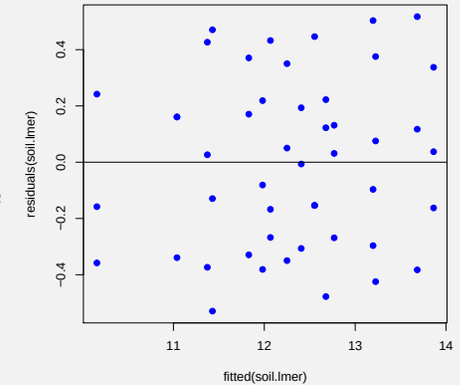
## Residual Plot

```

> plot(fitted(soil.lmer), residuals(soil.lmer))
> abline(h = 0)

```

- There are no bad patterns in the residual plot.

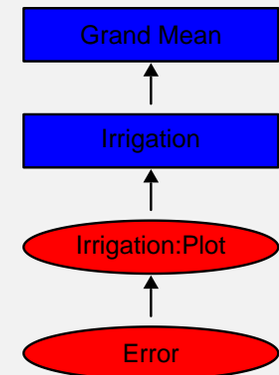


## Continue Analysis in R

- Draw a nesting diagram for the variables in the model.
- Use R to show alternative analysis using fixed effects.
- Draw connection to the correct ANOVA table calculation.

## Nesting

- A categorical variable  $A$  is *nested* in a categorical variable  $B$  if all of the observations within a single level of  $A$  are also within a single level of  $B$ .
- Irrigation method is nested within the grand mean.
- Plot is nested within irrigation method.
- Individual error is nested within each other variable.



## ANOVA

- For *balanced designs*, an appropriate  $F$ -test uses for its denominator the MS from *the closest nested random effect*.
- The correct p-value to test irrigation is only 0.0064, not  $10^{-14}$ .

```
> soil.lm = lm(moisture ~ irrigation + irrigation:plot)
> anova(soil.lm)

Analysis of Variance Table

Response: moisture
      Df Sum Sq Mean Sq F value    Pr(>F)
irrigation    3  29.4073    9.8024  76.507 1.097e-14 ***
irrigation:plot 12  17.3858    1.4488  11.308 2.342e-08 ***
Residuals    32   4.1000    0.1281
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

> anova(soil.lmer)

Analysis of Variance Table
      Df Sum Sq Mean Sq
irrigation  3  2.60086  0.86695

> c(9.8024/1.4488, 0.86695/0.1281)

[1] 6.765875 6.767760

> 1 - pf(6.76, 3, 12)

[1] 0.006385453
```