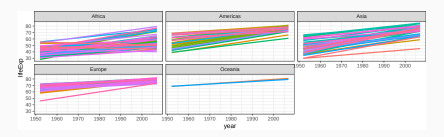
Multilevel models, part 2

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Interactions let us set a slope for each unit over time (a fixed effect growth model)

```
library(gapminder)
ggplot(gapminder, aes(x = year, y = lifeExp, color = country)) +
    geom_smooth(method = "lm", se=F) +
    guides(color = F) +
    facet_wrap(~continent)
```



Partially pooling slopes

- An interaction model presumes that all countries come from different populations
- · No information about the population of countries enters the model
- The unit-level interaction approach (fixed effects) constrains our modeling choices (time-invariant parameters can't be included)

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But, we can use the partial pooling technique to estimate a probability distribution for both intercepts *and* for slopes. We'll regularize the estimates, have parameters for the population we can use for inference, and have much more flexibility in our models.

Coffee shop wait times

- · Coffee shops have different wait times in the morning and afternoon
- Coffee shops with longer wait times in the morning will have a greater difference between morning and afternoon wait times (a floor effect)
- We want to pool information about coffee shops, presumably they are all similar in important ways
- Now, we'll pool information on both their intercepts (morning wait time), and their slopes (difference between morning and afternoon wait time)

Simulate coffee shop data

```
a<- 3.5
b<- -1
sigma_a<-1
sigma_b<-0.5
rho<- -0.7
```

Build a covariance matrix

These are symmetric matrices with variances on the diagonal, and covariances off-diagonal

So if our linear model is $y_i=\alpha_i+\beta_i x_i$ where α is a random slope and β is a random intercept, the covariance matrix can be written as

$$\left(\begin{array}{cc}\sigma_{\alpha}^{2} & \sigma_{\alpha}\sigma_{\beta}\rho_{\alpha\beta}\\ \sigma_{\alpha}\sigma_{\beta}\rho_{\alpha\beta} & \sigma_{\beta}^{2}\end{array}\right)$$

Compare to a correlation matrix

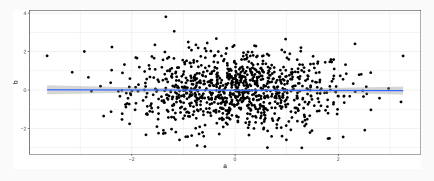
A correlation matrix represents correlations between random variables (covariances scaled to [-1, 1]).

$$\left(\begin{array}{cc} 1 & \rho_{\alpha\beta} \\ \rho_{\alpha\beta} & 1 \end{array}\right)$$

The impact of covariance on two random variables

Let's simulate a couple of examples: ho= 0, $\sigma_a=$ 1, $\sigma_b=$ 1

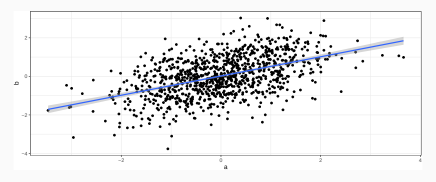
```
S<-matrix(c(1, 0, 0, 1), nrow=2)
mu<-c(0, 0)
sims<-mvrnorm(1000, mu, S)
plot_dat<-data.frame(a = sims[,1], b = sims[,2])
ggplot(plot_dat, aes(x = a, y = b)) +
    geom_point() +
    geom_smooth(method = "lm")</pre>
```



The impact of covariance on two random variables

Let's simulate a couple of examples: $\rho = 0.5, \sigma_a = 1, \sigma_b = 1$

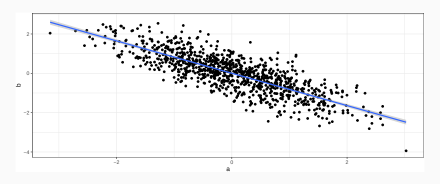
```
S<-matrix(c(1, 0.5, 0.5, 1), nrow=2)
mu<-c(0, 0)
sims<-mvrnorm(1000, mu, S)
plot_dat<-data.frame(a = sims[,1], b = sims[,2])
ggplot(plot_dat, aes(x = a, y = b)) +
    geom_point() +
    geom_smooth(method = "lm")</pre>
```



The impact of covariance on two random variables

Let's simulate a couple of examples: $ho = -0.8, \sigma_a = 1, \sigma_b = 1$

```
S<-matrix(c(1, -0.8, -0.8, 1), nrow=2)
mu<-c(0, 0)
sims<-mvrnorm(1000, mu, S)
plot_dat<-data.frame(a = sims[,1], b = sims[,2])
ggplot(plot_dat, aes(x = a, y = b)) +
geom_point() +
geom_smooth(method = "lm")</pre>
```



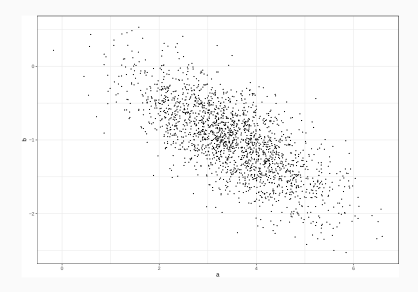
A covariance matrix can be obtained from standard deviations and correlations

```
sigmas<-c(sigma_a, sigma_b)
Rho<-matrix(c(1, rho, rho, 1), nrow=2)
# matrix multiplication produces \Sigma
Sigma<-diag(sigmas) %*% Rho %*% diag(sigmas)
Sigma

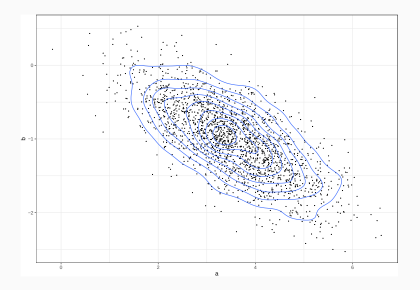
## [,1] [,2]
## [1,] 1.00 -0.35
## [2,] -0.35 0.25</pre>
```

Simulate the data

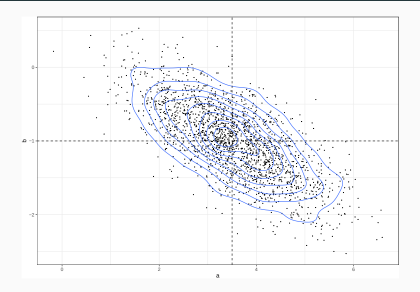
Plot the distribution



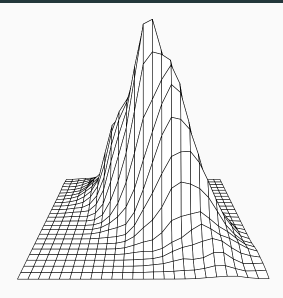
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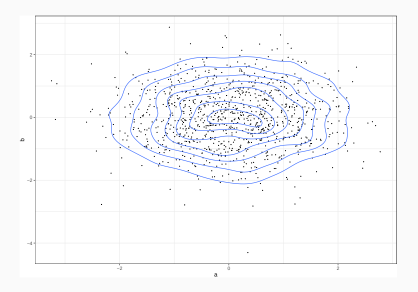
Plot the distribution



Seeing the contours in 3d



For comparison: no correlation between a and b



The setup

Our simulated data is the truth for 20 cafes. Now let's say we visit each of them 10 times to try to infer the truth.

```
N visits<-20
afternoon<-rep(0:1, N_visits * N_cafes/2)
cafe id<-rep(1:N cafes, each = N visits)
mu<-a cafe[cafe id] + b cafe[cafe id] * afternoon
sigma<-0.5
wait<-rnorm(N visits * N cafes, mu, sigma)</pre>
d<-data.frame(cafe = cafe id,</pre>
            afternoon = afternoon.
            wait = wait)
glimpse(d)
## Observations: 400
## Variables: 3
## $ cafe
             ## $ afternoon <int> 0, 1, 0, 1, 0, 1, 0, 1, 0, 1, 0, 1, 0, 1, 0, 1, 0, 1, 0, 1, 0, ...
## $ wait
            <dbl> 5.2069230, 2.5616650, 4.7259680, 2.2468053, 4.9995825, 3....
```

Defining the varying slopes model

Wait times (W) follow a normal likelihood, but each cafe can have it's own average wait time (intercept) and it's own average difference between morning and afternoon (A) waits (slope).

$$\begin{aligned} \textit{W}_{\textit{i}} &\sim \text{Normal}(\mu_{\textit{i}}, \sigma) \\ \mu_{\textit{i}} &= \alpha_{\text{cafe}_{\textit{i}}} + \beta_{\text{cafe}_{\textit{i}}} A_{\textit{i}} \\ \left[\begin{array}{c} \alpha_{\text{cafe}} \\ \beta_{\text{cafe}} \end{array}\right] &\sim \text{MVNormal}\left(\left[\begin{array}{c} \alpha \\ \beta \end{array}\right], S\right) \end{aligned}$$

The covariance matrix for an MVNormal variable

We can write a covariance matrix here as

$$S = \begin{pmatrix} \sigma_{\alpha}^2 & \sigma_{\alpha\beta} \\ \sigma_{\alpha\beta} & \sigma_{\beta}^2 \end{pmatrix}$$

Or write it as the product of the standard deviations (twice to square) and the correlation matrix R

$$S = \begin{pmatrix} \sigma_{\alpha} & 0 \\ 0 & \sigma_{\beta} \end{pmatrix} R \begin{pmatrix} \sigma_{\alpha} & 0 \\ 0 & \sigma_{\beta} \end{pmatrix}$$

The priors and hyper-priors for the varying intercepts and slopes

$$\alpha \sim \text{Normal}(5,2)$$

$$\beta \sim \text{Normal}(-1, 0.5)$$

$$\sigma \sim \text{Exponential}(1)$$

$$\sigma_{\alpha} \sim \text{Exponential}(1)$$

$$\sigma_{\beta} \sim \text{Exponential}(1)$$

$$R \sim \text{LKJcorr}(2)$$

A prior for a correlation matrix

A correlation matrix for two variables takes the form:

$$R = \begin{pmatrix} 1 & \rho \\ \rho & 1 \end{pmatrix}$$

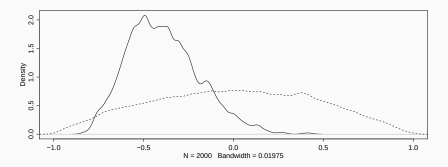
The LKJ correlation distribution samples correlation matrices where ρ can take on values on [-1, 1]. Higher values for the shape parameter makes the prior skeptical of extreme values (near -1 or 1).

Fitting the model with ulam()

```
m_cafe<-ulam(alist(
    wait ~ dnorm(mu, sigma),
    mu<-a_cafe[cafe] + b_cafe[cafe] * afternoon,
    c(a_cafe, b_cafe)[cafe] ~ multi_normal(c(a,b), Rho, sigma_cafe),
    a ~ dnorm(5, 2),
    b ~ dnorm(-1, 0.5),
    sigma_cafe ~ exponential(1),
    sigma ~ exponential(1),
    Rho ~ lkj_corr(2)
), data = d, chains = 4, cores = 4)</pre>
```

What the model tells us about ρ , prior and posterior

```
post <- extract.samples(m_cafe)
dens( post$Rho[,1,2] , xlim=c(-1,1) ) # posterior
R <- rlkjcorr( 1e4 , K=2 , eta=2 ) # prior
dens( R[,1,2] , add=TRUE , lty=2 )</pre>
```



Shrinks each posterior estimate toward group averages through pooling information

- Pools information on the correlation of slopes and intercepts (
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Visualizing adaptive regularization: set up plot

```
## compute observed a and b for each cafe
observed<-d %>%
  group by(cafe, afternoon) %>%
  summarise(a = mean(wait)) %>%
  pivot_wider(id_cols = cafe,
              values_from = a,
              names from = afternoon,
              names prefix = "a")
observed<-observed%>%
  mutate(b = a1 - a0) %>%
 ungroup()
## attach posterior means for a and b for each cafe
post<-extract.samples(m cafe)</pre>
post a<-apply(post$a cafe, 2, mean)
post_b<-apply(post$b_cafe, 2, mean)</pre>
plot dat<-observed %>%
  mutate(post a = post a, post b = post b)
```

The observed and posterior mean for α and β for each cafe



Let's return to the life expectancy data

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- · All countries saw life expectancy go up over time
- But rates of increase differ across places
- Countries that started with high life expectancy probably saw low growth, as there's a natural limit on how high it can go
- Countries with low life expectancy in the 1950s had much higher potential gains

Proposing a model

$$\begin{aligned} L_{i} &\sim \text{Normal}(\mu, \sigma) \\ \mu_{i} &= \alpha_{[\text{country}]i} + \beta_{[\text{country}]i} \times \textit{year}_{i} \\ \begin{bmatrix} \alpha_{\text{country}} \\ \beta_{\text{country}} \end{bmatrix} &\sim \text{MVNormal} \begin{pmatrix} \begin{bmatrix} \alpha \\ \beta \end{bmatrix}, S \end{pmatrix} \\ S &= \begin{pmatrix} \sigma_{\alpha} & 0 \\ 0 & \sigma_{\beta} \end{pmatrix} R \begin{pmatrix} \sigma_{\alpha} & 0 \\ 0 & \sigma_{\beta} \end{pmatrix} \\ &\alpha &\sim \text{Normal}(0, 2) \\ &\beta &\sim \text{Normal}(0, 2) \\ &\sigma, \sigma_{\alpha}, \sigma_{\beta} &\sim \text{Exponential}(1) \end{aligned}$$

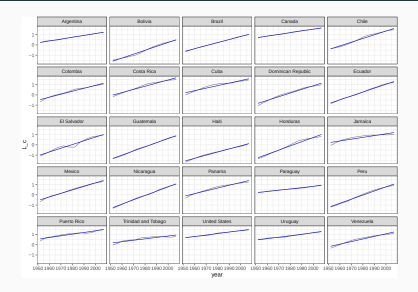
 $R \sim LKJcorr(2)$

30

Estimating the model

```
m_gm1<-ulam(alist(
  L_c ~ dnorm(mu, sigma),
  mu <- alpha[country] + beta[country] * year_c,
  c(alpha, beta)[country] ~ multi_normal(c(0, b), Rho, sigma_country
  b ~ dnorm(0, 2),
  sigma_country ~ dexp(1),
  sigma ~ dexp(1),
  Rho ~ lkj_corr(1)
), data = dat, chains = 4, cores = 4)</pre>
```

Visualizing the inferences

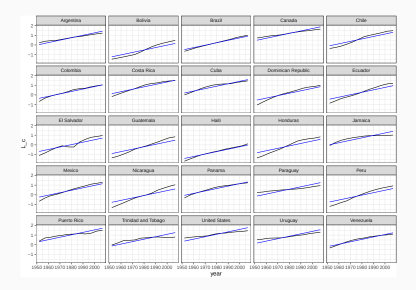


Compare to alternative models

Let's compare this to a model with a single slope and variable intercepts

```
m_gm2<-ulam(alist(
    L_c ~ dnorm(mu, sigma),
    mu <- alpha[country] * beta * year_c,
    alpha[country] ~ dnorm(a, sigma_country),
    a ~ dnorm(0,2),
    beta ~ dnorm(0, 2),
    sigma_country ~ dexp(1),
    sigma ~ dexp(1)
), data = dat, chains = 4, cores = 4)</pre>
```

Visualize the results



What did we learn about country parameters

```
precis(m_gm1, pars = "sigma_country", depth = 3)
```

```
## mean sd 5.5% 94.5% n_eff Rhat4
## sigma_country[1] 1.18160832 0.070861593 1.0708565 1.29840568 4547.179 0.9985532
## sigma_country[2] 0.06481852 0.004645245 0.0578517 0.07257805 2791.211 0.9997337
```

Summary

- We can use partial pooling for slopes just as we did for intercepts
- This treats both the intercepts and slopes as coming from the same population
- This makes a lot of sense when our units are exchangeable, or come from the same population
- Make sure to read the chapter for some advanced examples, including applications for causal inference
- Homework: Ch14 easy and medium questions, H1 if you want a challenge