Bayesian Statistics with Stan

Packages for this section

```
library(tidyverse)
# library(rstan)
library(cmdstanr)
library(posterior)
library(bayesplot)
```

Installation 1/2

o cmdstanr:

• posterior and bayesplot, from the same place:

Installation 2/2

Then, to check that you have the C++ stuff needed to compile Stan code: $check_cmdstan_toolchain()$

The C++ toolchain required for CmdStan is setup properly!
and then:

```
install cmdstan(cores = 2)
```

If you happen to know how many cores (processors) your computer has, insert the appropriate number. (My laptop has 4 and my desktop 6.)

All of this is done once. If you have problems, go here (link).

Bayesian and frequentist inference

- The inference philosophy that we have learned so far says that:
 - parameters to be estimated are *fixed* but *unknown*
 - Data random; if we took another sample we'd get different data.
- This is called "frequentist" or "repeated-sampling" inference.
- Bayesian inference says:
 - parameters are random, data is given
- Ingredients:
 - **prior distribution**: distribution of parameters before seeing data.
 - likelihood: model for data if the parameters are known
 - posterior distribution: distribution of parameters after seeing data.

Distribution of parameters

- Instead of having a point or interval estimate of a parameter, we have an entire distribution
- so in Bayesian statistics we can talk about eg.
 - probability that a parameter is bigger than some value
 - probability that a parameter is close to some value
 - probability that one parameter is bigger than another
- Name comes from Bayes' Theorem, which here says posterior is proportional to likelihood times prior
- more discussion about this is in a blog post.

An example

• Suppose we have these (integer) observations:

```
(x \leftarrow c(0, 4, 3, 6, 3, 3, 2, 4))
```

[1] 0 4 3 6 3 3 2 4

- ullet Suppose we believe that these come from a Poisson distribution with a mean λ that we want to estimate.
- We need a prior distribution for λ . I will (for some reason) take a Weibull distribution with parameters 1.1 and 6, that has quartiles 2 and 6. Normally this would come from your knowledge of the data-generating *process*.
- The Poisson likelihood can be written down (see over).

Some algebra

 \bullet We have n=8 observations $x_i,$ so the Poisson likelihood is proportional to

$$\prod_{i=1}^{n} e^{-\lambda} \lambda^{x_i} = e^{-n\lambda} \lambda^S,$$

where $S = \sum_{i=1}^{n} x_i$.

• then you write the Weibull prior density (as a function of λ):

$$C(\lambda/6)^{0.1}e^{-(\lambda/6)^{1.1}}$$

where C is a constant.

 and then you multiply these together and try to recognize the distributional form. Only, here you can't. The powers 0.1 and 1.1 get in the way.

Sampling from the posterior distribution

- Wouldn't it be nice if we could just sample from the posterior distribution? Then we would be able to compute it as accurately as we want.
- Metropolis and Hastings: devise a Markov chain (C62) whose limiting distribution is the posterior you want, and then sample from that Markov chain (easy), allowing enough time to get close enough to the limiting distribution.
- Stan: uses a modern variant that is more efficient (called Hamiltonian Monte Carlo), implemented in R packages cmdstanr.
- Write Stan code in a file, compile it and sample from it.

Components of Stan code: the model

```
model {
    // likelihood
    x ~ poisson(lambda);
}
```

This is how you say "X has a Poisson distribution with mean λ ". Note that lines of Stan code have semicolons on the end.

Components of Stan code: the prior distribution

```
model {
   // prior
   lambda ~ weibull(1.1, 6);
   // likelihood
   x ~ poisson(lambda);
}
```

Components of Stan code: data and parameters

```
• first in the Stan code:
data {
   int x[8];
}
parameters {
   real<lower=0> lambda;
}
```

Compile and sample from the model

compile

```
poisson1 <- cmdstan_model("poisson1.stan")</pre>
  set up data
```

poisson1 data \leftarrow list(x = x)

```
sample
```

```
## Running MCMC with 4 sequential chains...
```

poisson1_fit <- poisson1\$sample(data = poisson1_data)</pre>

##

Chain 1 Iteration: Chain 1 Iteration:

Chain 1 Iteration:

(11-----)

100 / 2000 [5%]

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The output

```
poisson1_fit
```

```
## variable mean median sd mad q5 q95 rhat
## lp_ 3.75 4.03 0.71 0.31 2.36 4.26 1.00
## lambda 3.19 3.15 0.63 0.61 2.24 4.31 1.00
## ess_bulk ess_tail
## 1827 2086
## 1443 1807
```

Comments

- ullet This summarizes the posterior distribution of λ
- the posterior mean is 3.19
- with a 90% posterior interval of 2.24 to 4.31.
- The probability that λ is between these two values really is 90%.

Making the code more general

• The coder in you is probably offended by hard-coding the sample size and the parameters of the prior distribution. More generally:

```
data {
  int<lower=1> n;
  real<lower=0> a;
  real<lower=0> b;
  int x[n];
model {
// prior
lambda ~ weibull(a, b);
// likelihood
x ~ poisson(lambda);
}
```

Set up again and sample:

Compile again:

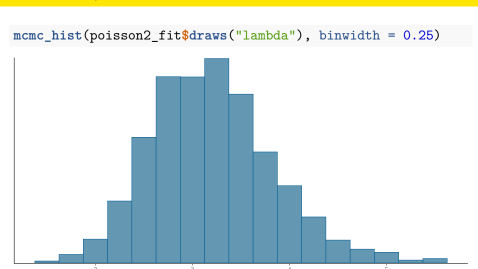
```
poisson2 <- cmdstan_model("poisson2.stan")</pre>
poisson2
## // Estimating Poisson mean
##
## data {
##
     int<lower=1> n:
## real<lower=0> a;
## real<lower=0> b:
## int x[n]:
## }
##
## parameters {
     real<lower=0> lambda;
##
## }
```

output should be the same (to within randomness)

```
poisson2_fit
## variable mean median sd mad q5 q95 rhat
```

```
## lp_ 3.76 4.03 0.72 0.31 2.31 4.26 1.00
## lambda 3.21 3.16 0.63 0.60 2.29 4.30 1.00
## ess_bulk ess_tail
## 1707 1844
## 1429 1830
```

Picture of posterior



lambda

Extracting actual sampled values

```
A little awkward at first:
poisson2 fit$draws()
```

##

```
## # A draws_array: 1000 iterations, 4 chains, and 2 variables
## , , variable = lp__
##
##
           chain
## iteration 1 2 3 4
          1 4.2 3.6 3.9 4.0
##
          2 4.2 3.5 4.1 4.0
##
##
          3 3.9 3.8 4.1 3.6
##
          4 4.0 3.1 3.8 4.2
          5 4.0 4.0 4.0 4.2
##
##
    , variable = lambda
##
```

Sampled values as dataframe

as_draws_df(poisson2_fit\$draws()) %>%

```
as_tibble() -> poisson2_draws
poisson2_draws
## # A tibble: 4,000 x 5
##
      lp lambda .chain .iteration .draw
     <dbl> <dbl> <int>
##
                             <int> <int>
##
  1 4.20 2.99
##
   2 4.20 2.99
   3 3.93 3.72
##
   4 4.04 3.62
##
                                 4
##
   5 4.02 3.64
                                 5
                                      5
##
   6 3.74 3.87
                                 6
##
   7 1.70 4.82
##
   8 4.07 2.83
                                 8
     4.16 3.47
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```

Posterior predictive distribution

 Another use for the actual sampled values is to see what kind of response values we might get in the future. This should look something like our data. For a Poisson distribution, the response values are integers:

```
poisson2_draws %>%
  rowwise() %>%
  mutate(xsim = rpois(1, lambda)) -> d
d
```

```
## # A tibble: 4,000 x 6

## lp_ lambda .chain .iteration .draw xsim

## <dbl> <dbl> <int> <int> <int> <int> <int> <int> <int> <int> <int> <int > < <int > <
```

Comparison

Our actual data values were these:

```
x
```

```
## [1] 0 4 3 6 3 3 2 4
```

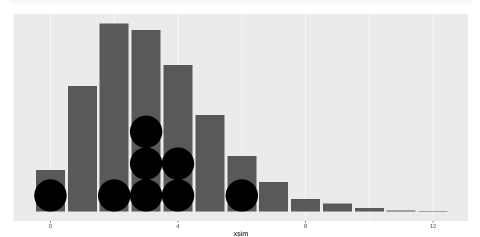
- None of these are very unlikely according to our posterior predictive distribution, so our model is believable.
- Or make a plot: a bar chart with the data on it as well (over):

```
ggplot(d, aes(x = xsim)) + geom_bar() +
geom_dotplot(data = tibble(x), aes(x = x), binwidth = 1) +
scale_y_continuous(NULL, breaks = NULL) -> g
```

• This also shows that the distribution of the data conforms well enough to the posterior predictive distribution (over).

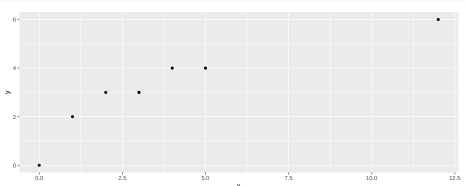
The plot

g



Do they have the same distribution?

```
qqplot(d$xsim, x, plot.it = F) %>% as_tibble() -> dd
ggplot(dd, aes(x=x, y=y)) + geom_point()
```



the observed zero is a bit too small compared to expected (from the posterior), but the other points seem pretty well on a line.

Analysis of variance, the Bayesian way

Recall the jumping rats data:

```
my url <-
  "http://ritsokiguess.site/datafiles/jumping.txt"
rats0 <- read_delim(my_url, " ")
rats0
## # A tibble: 30 x 2
##
   group density
##
     <chr> <dbl>
```

```
611
##
   1 Control
   2 Control 621
##
   3 Control 614
##
##
   4 Control
                 593
##
   5 Control
                 593
##
   6 Control
                 653
   7 Control
                 600
##
```

Our aims here

- Estimate the mean bone density of all rats under each of the experimental conditions
- \bullet Model: given the group means, each observation normally distributed with common variance σ^2
- Three parameters to estimate, plus the common variance.
- Obtain posterior distributions for the group means.
- Ask whether the posterior distributions of these means are sufficiently different.

Numbering the groups

rats0 %>% mutate(

Stan doesn't handle categorical variables (everything is real or int).

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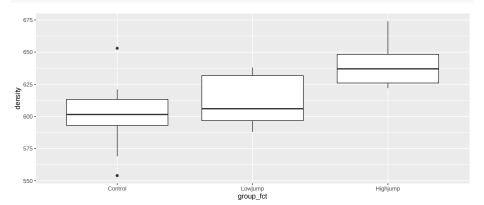
- Turn the groups into group *numbers* first.
- Take opportunity to put groups in logical order:

```
group_fct = fct_inorder(group),
 group_no = as.integer(group_fct)
) -> rats
rats
## # A tibble: 30 \times 4
##
     group density group_fct group_no
##
     <chr> <dbl> <fct> <int>
##
   1 Control 611 Control
##
   2 Control 621 Control
##
   3 Control 614 Control
   4 Control 593 Control
##
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```

Plotting the data 1/2

Most obviously, boxplots:

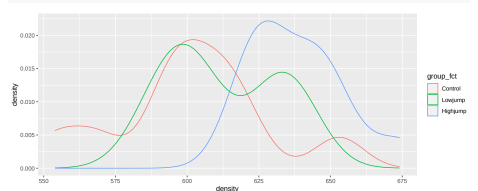
```
ggplot(rats, aes(x = group_fct, y = density)) +
  geom_boxplot()
```



Plotting the data 2/2

Another way: density plot (smoothed out histogram); can distinguish groups by colours:

```
ggplot(rats, aes(x = density, colour = group_fct)) +
  geom_density()
```



The procedure

- For each observation, find out which (numeric) group it belongs to,
- then model it as having a normal distribution with that group's mean and the common variance.
- Stan does for loops.

The model part

```
Suppose we have n_obs observations:

model {
    // likelihood
    for (i in 1:n_obs) {
        g=group_no[i];
        density[i] ~ normal(mu[g], sigma);
    }
}
```

The variables here

- on obs is data.
- g is a temporary integer variable only used here
- i is only used in the loop (integer) and does not need to be declared
- density is data, a real vector of length n_obs
- mu is a parameter, a real vector of length 3 (3 groups)
- sigma is a real parameter

mu and sigma need prior distributions:

- for mu, each component independently normal with mean 600 and SD
 50 (my guess at how big and variable they will be)
- for sigma, chi-squared with 50 df (my guess at typical amount of variability from obs to obs)

Complete the model section:

```
model {
  int g;
  // priors
  mu ~ normal(600, 50);
  sigma ~ chi_square(50);
  // likelihood
  for (i in 1:n obs) {
    g=group_no[i];
    density[i] ~ normal(mu[g], sigma);
```

Parameters

The elements of mu, one per group, and also sigma, scalar, lower limit zero:

```
parameters {
  real mu[n_group];
  real<lower=0> sigma;
}
```

- Declare sigma to have lower limit zero here, so that the sampling runs smoothly.
- declare n_group in data section

Data

```
Everything else:
data {
  int n_obs;
  int n_group;
  real density[n_obs];
  int<lower=1, upper=n_group> group_no[n_obs];
}
```

Compile

Arrange these in order data, parameters, model in file anova.stan, then:

anova <- cmdstan_model("anova.stan")</pre>

Set up data and sample

Chain 1 Tteration:

```
Supply values for everything declared in data:
anova data <- list(
  n_{obs} = 30,
  n_{group} = 3,
  density = rats$density,
  group_no = rats$group_no
anova fit <- anova$sample(data = anova data)
## Running MCMC with 4 sequential chains...
##
## Chain 1 Iteration:
                          1 / 2000 [ 0%]
                                            (Warmup)
                                            (Warmup)
## Chain 1 Iteration: 100 / 2000 [ 5%]
                                            (Warmup)
## Chain 1 Iteration: 200 / 2000 [ 10%]
                                            (Warmup)
## Chain 1 Iteration: 300 / 2000 [ 15%]
```

/ 2000 [20%]

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(Warmup)

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Check that the sampling worked properly

```
anova fit$cmdstan diagnose()
## Processing csv files: /tmp/Rtmpcpl9DP/anova-202107061207-1-4bf7a0.csv, /tmp/Rtmp
202107061207-2-4bf7a0.csv, /tmp/Rtmpcpl9DP/anova-202107061207-3-4bf7a0.csv, /tmp/Rt
202107061207-4-4bf7a0.csv
##
## Checking sampler transitions treedepth.
## Treedepth satisfactory for all transitions.
##
## Checking sampler transitions for divergences.
## No divergent transitions found.
##
## Checking E-BFMI - sampler transitions HMC potential energy.
## E-BFMI satisfactory.
##
## Effective sample size satisfactory.
##
## Split R-hat values satisfactory all parameters.
##
## Processing complete, no problems detected.
```

Look at the results

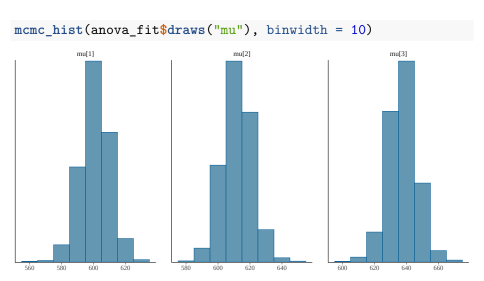
anova_fit

```
variable mean median sd mad q5 q95
##
##
      lp_ -41.02 -40.67 1.47 1.29 -43.77 -39.29
##
      mu[1] 600.99 600.95 8.92 8.68 586.34 615.31
##
      mu[2] 612.09 612.14 9.15 9.21 597.36 627.05
##
      mu[3] 637.45 637.48 8.99 8.75 622.83 652.12
      sigma 28.69 28.35 4.15 4.07 22.57 36.17
##
##
   rhat ess bulk ess tail
##
   1.00
           1707
                   2281
##
   1.00 3644 2708
##
   1.00 3892 2334
  1.00 4366 2746
##
  1.00 2713
##
                   2619
```

Comments

- The posterior 95% intervals for control (group 1) and highjump (group 3) do not quite overlap, suggesting that these exercise groups really are different.
- Bayesian approach does not normally do tests: look at posterior distributions and decide whether they are different enough to be worth treating as different.

Plotting the posterior distributions for the mu



Extract the sampled values

```
as_draws_df(anova_fit$draws()) %>% as_tibble() -> anova_draws
anova_draws
## # A tibble: 4,000 x 8
     lp__ `mu[1]` `mu[2]` `mu[3]` sigma .chain
##
## <dbl> <dbl> <dbl> <dbl> <int>
## 1 -43.7 599. 588. 642. 26.0
## 2 -46.0 618. 640. 626. 36.7
  3 -41.7 596. 594. 642. 26.8
##
  4 -42.0 606. 632. 631. 28.8
##
##
  5 -40.2 600. 602. 635. 31.1
```

6 -45.1 611. 635. 656. 29.4 ## 7 -41.1 598. 622. 650. 31.9 ## 8 -41.3 597. 600. 649. 32.2 ## 9 -40.1 595. 603. 643. 29.5 ## 10 -39.6 603. 619. 631. 25.5

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estimated probability that $\mu_3>\mu_1$

```
anova_draws %>%
  count(`mu[3]`>`mu[1]`) %>%
  mutate(prob = n/sum(n))
```

High jumping group almost certainly has larger mean than control group.

More organizing

- for another plot
 - make longer
 - give group values their proper names back

What we have now:

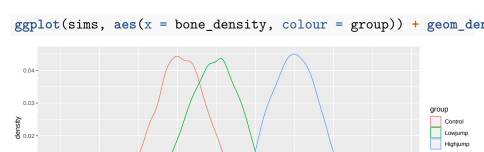
sims %>% sample n(8)

```
## # A tibble: 8 x 7
##
     lp__ sigma .chain .iteration .draw group
## <dbl> <dbl> <int>
                          <int> <int> <fct>
## 1 -40.4 24.3
                            173 173 Highjump
                            951 951 Lowjump
## 2 -40.8 28.1
## 3 -40.3 31.7
                         209 209 Control
## 4 -39.7 25.6
                    3
                         712 2712 Highjump
## 5 -41.0 33.1
                           998
                                 998 Lowjump
                         728 2728 Lowjump
## 6 -39.7 29.1
                    3
## 7 -40.8 29.9
                    4
                         709 3709 Highjump
## 8 -41.6 32.1
                    3
                            328
                                2328 Lowjump
## # ... with 1 more variable: bone density <dbl>
```

Density plots of posterior mean distributions

0.01 -

0.00 -



bone density

Posterior predictive distributions

A tibble: 12,000 x 8

Randomly sample from posterior means and SDs in sims. There are 12000 rows in sims:

```
sims %>% mutate(sim_data = rnorm(12000, bone_density, sigma))
ppd
```

```
##
       lp__ sigma .chain .iteration .draw group
##
      <dbl> <dbl> <int>
                              <int> <int> <fct>
## 1 -43.7 26.0
                                          1 Control
   2 -43.7 26.0
##
                                          1 Lowjump
##
   3 -43.7 26.0
                                          1 Highjump
##
   4 -46.0 36.7
                                          2 Control
   5 -46.0 36.7
##
                                          2 Lowjump
                                          2 Highjump
##
   6 -46.0 36.7
## 7 -41.7 26.8
                                   3
                                          3 Control
##
    8 -41.7 26.8
                                          3 Lowjump
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```

Compare posterior predictive distribution with actual data

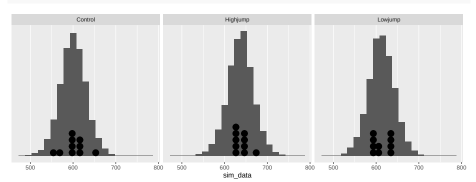
- Check that the model works: distributions of data similar to what we'd predict
- Idea: make plots of posterior predictive distribution, and plot actual data as points on them
- Use facets, one for each treatment group:

```
my_binwidth <- 15
ggplot(ppd, aes(x = sim_data)) +
    geom_histogram(binwidth = my_binwidth) +
    geom_dotplot(
        data = rats, aes(x = density),
        binwidth = my_binwidth
) +
    facet_wrap(~group) +
    scale_y_continuous(NULL, breaks = NULL) -> g
```

- See (over) that the data values are mainly in the middle of the predictive distributions.
- Even for the control group that had outliers.

The plot





Extensions

- if you want a different model other than normal, change distribution in model section
- if you want to allow unequal spreads, create sigma[n_group] and in model density[i] ~ normal(mu[g], sigma[g]);
- Stan will work just fine after you recompile
- very flexible.
- Typical modelling strategy: start simple, add complexity as warranted by data.