# **Assignment 7**

## Hierarchical model in Stan

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### 1 General information

This is the template for assignment 7. You can download the separate model with bad priors and the qmd-file or copy the code from this rendered document after clicking on </> Code in the top right corner.

Please replace the instructions in this template by your own text, explaining what you are doing in each exercise.

## 2 Hierarchical Model: Chicken Data with Stan (6p)

## 2.1 Choosing a weakly informative prior by intuition

- 2.1.1 (a)
- 2.1.2 (b)
- 2.1.3 (c)
- 2.1.4 (d)
- 2.1.5 (e)

### 2.2 Choosing a weakly informative prior using external references

- 2.2.1 (f)
- 2.2.2 (g)
- 2.2.3 (h)
- 2.2.4 (i)

### 2.3 Non-normal priors

2.3.1 (j)

## 2.4 Modeling diet effects on chicken weight

```
Data inside, don't peek before you have set your priors!
 Have you set your priors?
    data("ChickWeight")
    Chick12 <- ChickWeight |> filter(Time == 12)
    head(Chick12)
 Grouped Data: weight ~ Time | Chick
   weight Time Chick Diet
       106
             12
                     1
                          1
       122
 2
             12
                     2
                          1
 3
       115
             12
                     3
                          1
 4
       102
             12
                     4
                          1
 5
       141
             12
                     5
                          1
       141
             12
                     6
```

### 2.4.2 (I)

For the figures below, we use the earlier samples for the separate model with bad priors. When you have implemented the pooled and hierarchical models, edit the code below to include samples from your model into the figures.

### 2.4.3 (m)

## Mean of diet 4

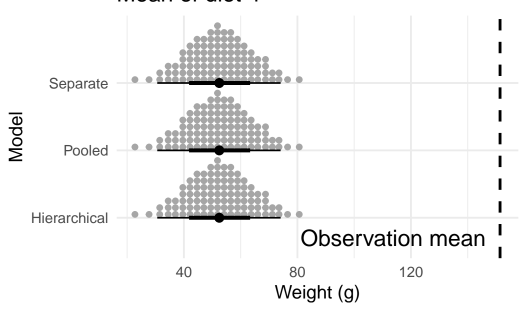


Figure 1: Posterior distribution of the mean weight of chicks consuming diet 4.

### 2.4.4 (n)

```
ggplot(predicted_weight_diet_4, aes(x = predicted_weight, y = model_name)) +
   stat_dotsinterval(quantiles = 100, scale = .9) +
   vline_at(diet_means[4], size = 1, linetype = "dashed") +
   # Annotate the vline from above.
```

## Weigth of a chick with diet 4

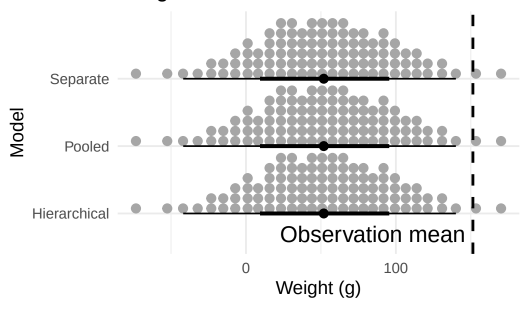


Figure 2: The (posterior) predictive distribution of the weight of a chick consuming diet 4.

### 2.4.5 (o)

## Mean of a new diet

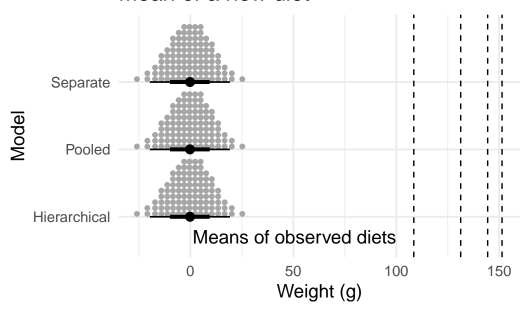


Figure 3: Posterior distribution of the mean weight of chicks consuming the new diet 5 not seen before.

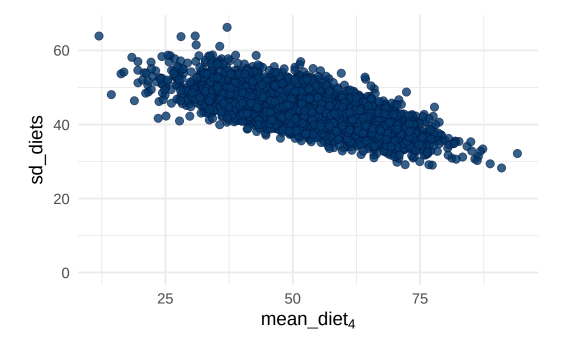
### 2.4.6 (p)

## 3 Hierarchical model with BRMS (3p)

### 3.0.1 (a)

Scale for y is already present.

Adding another scale for y, which will replace the existing scale.



### 3.0.2 (b)

Because brms is a bit chatty, suppress its output in the PDF using the block above, but copy the code you executed into the code block below, which doesn't execute:

```
# Copy the you used to fit the brms model here
```

### 3.0.3 (c)

```
# Draws for mu_4
mu_4 = posterior_epred(brms_fit, newdata = data.frame(Diet=4))

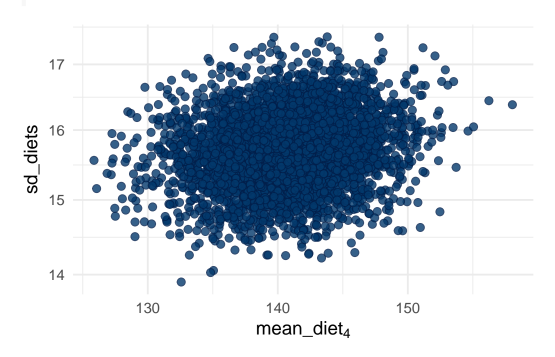
# Compute the mean, and quantiles. Remember to round your answers accordingly.
# ...
```

### 3.0.4 (d)

#### 3.0.5 Scatterplot for non-centered parametrization

Due the non-centered parametrization, we need to transform compute the  $\mu_d$  term as the sum of the population intercept and the group specific deviation from the intercept. You can choose which diet to plot by modifying the d integer in r\_Diet[d,Intercept].

```
scale_y_log10() +
xlab(expression(mean_diet[4])) +
ylab(expression(sd_diets))
```



3.0.6 (e)